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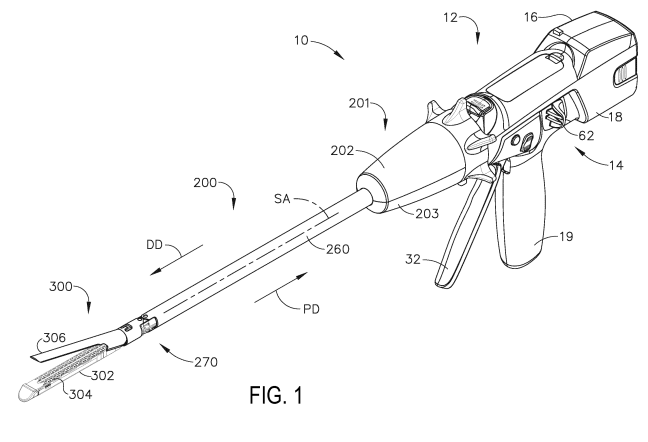
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(54) MODULAR SURGICAL INSTRUMENT WITH CONFIGURABLE OPERATING MODE

(57) Various examples are directed to systems and methods for operating and controlling a modular surgical instrument comprising an operating mode. A controller may receive electrical signal associated with an activa-

tion of at least one user-actuated input device electrically coupled to the controller and override the operating mode of the surgical instrument upon receiving the electrical signal.



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Description

FIELD

[0001] The present disclosure relates to surgical instruments and, in various circumstances, to surgical stapling and cutting instruments and staple cartridges therefor that are designed to staple and cut tissue.

BACKGROUND

[0002] In a motorized surgical stapling and cutting instrument, it would be useful to provide a user actuatable control which overrides any limitation imposed by the instrument. Furthermore, it would be useful to automatically drive the cutting member drive bar to target positions at a target velocity in order to configure internal mechanisms. It would be useful to provide this functionality to automatically target positions and to minimize user concern and operating time. While several devices have been made and used, it is believed that no one prior to the inventors has made or used the device described in the appended claims.

BRIEF SUMMARY

[0003] In one aspect of the invention, a surgical instrument is provided. The surgical instrument, comprising: a drive system; an electric motor mechanically coupled to the drive system; a controller electrically coupled to the electric motor, and at least one user-actuated input device electrically coupled to the controller; and wherein the controller is configured to operate the electric motor; wherein the controller is configured to detect activation of the at least one user-actuated input device and to override the operating mode for the electric motor upon detecting the activation of the at least one user-actuated input device.

[0004] In another aspect of the invention, a surgical instrument is provided. The surgical instrument, comprising: a drive system; an end effector comprising an anvil, an elongated channel mechanically coupled to the anvil, a staple cartridge supported by the elongated channel and a cutting member mechanically coupled to the drive system; an electric motor mechanically coupled to the drive system; a controller electrically connected to the electric motor; a first sensor electrically coupled to the controller and configured to sense a first parameter associated with the anvil; a second sensor electrically coupled to the controller and configured to sense a second parameter associated with the cutting member; and at least one user-actuated input device electrically coupled to the controller; and wherein the controller is configured to implement an operating mode for the electric motor based on at least one of the first parameter and the second parameter; and wherein the controller is configured such that activation of the at least one user-actuated input

device causes the controller to override the operating mode for the electric motor.

[0005] The foregoing summary is illustrative only and is not intended to be in any way limiting. In addition to the illustrative aspects and features described above, further aspects and features will become apparent by ref-

erence to the drawings and the following detailed description.

10 BRIEF DESCRIPTION OF THE DRAWINGS

[0006] The novel features of the aspects described herein are set forth with particularity in the appended claims. The aspects, however, both as to organization and methods of operation may be better understood by reference to the following description, taken in conjunction with the accompanying drawings as follows.

FIG. 1 is a perspective view of a surgical instrument that has an interchangeable shaft assembly operably coupled thereto in accordance with one or more aspects of the present disclosure.

FIG. 2 is an exploded assembly view of the interchangeable shaft assembly and surgical instrument of FIG. 1 in accordance with one or more aspects of the present disclosure.

FIG. 3 is another exploded assembly view showing portions of the interchangeable shaft assembly and surgical instrument of FIGS. 1 and 2 in accordance with one or more aspects of the present disclosure.
FIG. 4 is an exploded assembly view of a portion of the surgical instrument of FIGS. 1-3 in accordance with one or more aspects of the present disclosure.
FIG. 5 is a cross-sectional side view of a portion of the surgical instrument of FIG. 4 with the firing trigger in a fully actuated position in accordance with one or more aspects of the present disclosure.

FIG. 6 is another cross-sectional view of a portion of the surgical instrument of FIG. 5 with the firing trigger in an unactuated position in accordance with one or more aspects of the present disclosure.

FIG. 7 is another exploded assembly view of portions of the interchangeable shaft assembly of FIG. 7 in accordance with one or more aspects of the present disclosure.

FIG. 8 is a cross-sectional view of a portion of the interchangeable shaft assembly of FIGS. 7-9, in accordance with one or more aspects of the present disclosure.

FIG. 9 is another perspective view of the portion of an interchangeable shaft assembly with the switch drum mounted thereon in accordance with one or more aspects of the present disclosure.

FIG. 10 is a perspective view of a portion of the interchangeable shaft assembly of FIG. 11 operably coupled to a portion of the surgical instrument of FIG.1 illustrated with the closure trigger thereof in an unactuated position in accordance with one or more

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aspects of the present disclosure.

FIG. 11 is a right side elevational view of the interchangeable shaft assembly and surgical instrument of FIG.10 in accordance with one or more aspects of the present disclosure.

FIG. 12 is a perspective view of a portion of the interchangeable shaft assembly of FIG. 11 operably coupled to a portion of the surgical instrument of FIG. 1 illustrated with the closure trigger thereof in an actuated position and a firing trigger thereof in an unactuated position in accordance with one or more aspects of the present disclosure.

FIG. 13 is a right side elevational view of the interchangeable shaft assembly operably coupled to a portion of the surgical instrument of FIG. 1 illustrated with the closure trigger thereof in an actuated position and the firing trigger thereof in an actuated position in accordance with one or more aspects of the present disclosure.

FIG. 14 is an exploded view of one aspect of an end ²⁰ effector of the surgical instrument of FIG. 1 in accordance with one or more aspects of the present disclosure.

FIG. 15 is a schematic of a system for powering down an electrical connector of a surgical instrument handle when a shaft assembly is not coupled thereto in accordance with one or more aspects of the present disclosure.

FIGS. 16A-16B is a circuit diagram of the surgical instrument of FIG. 1 spanning two drawings sheets in accordance with one or more aspects of the present disclosure

FIGS. 17A-17B, is a circuit diagram of the surgical instrument of FIG. 1 in accordance with one or more aspects of the present disclosure.

FIG. 18 is a block diagram the surgical instrument of FIG. 1 illustrating interfaces between the handle assembly and the power assembly and between the handle assembly and the interchangeable shaft assembly in accordance with one or more aspects of the present disclosure.

FIG. 19 illustrates a logic diagram of a system for evaluating sharpness of a cutting edge of a surgical instrument in accordance with one or more aspects of the present disclosure.

FIG. 20 illustrates a logic diagram of a system for determining the forces applied against a cutting edge of a surgical instrument by a sharpness testing member at various sharpness levels in accordance with one or more aspects of the present disclosure.

FIG. 21 illustrates one aspect of a process for adapting operations of a surgical instrument in accordance with one or more aspects of the present disclosure. FIG. 22A depicts an example end-effector of a medical device surrounding tissue in accordance with one or more aspects of the present disclosure.

FIG. 22B depicts an example end-effector of a medical device compressing tissue in accordance with one or more aspects of the present disclosure. FIG. 23A depicts example forces exerted by an endeffector of a medical device compressing tissue in accordance with one or more aspects of the present disclosure.

FIG. 23B also depicts example forces exerted by an end-effector of a medical device compressing tissue in accordance with one or more aspects of the present disclosure.

FIG. 24 depicts an example tissue compression sensor system in accordance with one or more aspects of the present disclosure.

FIG. 25 also depicts an example tissue compression sensor system in accordance with one or more aspects of the present disclosure.

FIG. 26 also depicts an example tissue compression sensor system in accordance with one or more aspects of the present disclosure.

FIG. 27 depicts an example end-effector channel frame in accordance with one or more aspects of the present disclosure.

FIG. 28 depicts an example end-effector in accordance with one or more aspects of the present disclosure.

FIG. 29 also depicts an example end-effector channel frame in accordance with one or more aspects of the present disclosure.

FIG. 30 also depicts an example end-effector channel frame in accordance with one or more aspects of the present disclosure.

FIG. 31 also depicts an example end-effector channel frame in accordance with one or more aspects of the present disclosure.

FIG. 32 depicts an example electrode in accordance with one or more aspects of the present disclosure. FIG. 33 depicts an example electrode wiring system in accordance with one or more aspects of the present disclosure.

FIG. 34 also depicts an example end-effector channel frame in accordance with one or more aspects of the present disclosure.

FIG. 35 is an example circuit diagram in accordance with one or more aspects of the present disclosure. FIG. 36 is also an example circuit diagram in accord-

ance with one or more aspects of the present disclosure.

FIG. 37 is also an example circuit diagram in accordance with one or more aspects of the present disclosure.

FIG. 38 is a perspective view of a surgical instrument with an articulable, interchangeable shaft in accordance with one or more aspects of the present disclosure.

FIG. 39 is a side view of the tip of the surgical instrument shown in FIG. 38 in accordance with one or more aspects of the present disclosure.

FIG. 40 illustrates a cross-sectional view of an end effector of a surgical instrument in accordance with

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one or more aspects of the present disclosure.

FIG. 41 illustrates a logic diagram of a feedback system in accordance with one or more aspects of the present disclosure.

FIG. 42 illustrates a logic diagram of a feedback system in accordance with one or more aspects of the present disclosure.

FIG. 43 is a diagram of a smart sensor component in accordance with an aspect the present disclosure. FIG. 44 illustrates one aspect of a circuit configured to convert signals from a first sensor and a plurality of secondary sensors into digital signals receivable by a processor in accordance with one or more aspects of the present disclosure.

FIG. 45 illustrates one aspect of an exploded view of a staple cartridge that comprises a flex cable connected to a magnetic field sensor and processor in accordance with one or more aspects of the present disclosure.

FIG. 46 illustrates the end effector shown in FIG. 46 with a flex cable and without the shaft assembly in accordance with one or more aspects of the present disclosure.

FIGS. 47 and 48 illustrate an elongated channel portion of an end effector without the anvil or the staple cartridge, to illustrate how the flex cable shown in FIG. 46 can be seated within the elongated channel in accordance with one or more aspects of the present disclosure.

FIG. 49 illustrates a flex cable, shown in FIGS. 46-48, alone in accordance with one or more aspects of the present disclosure.

FIG. 50 illustrates a close up view of the elongated channel shown in FIG. 114 and 115 with a staple cartridge coupled thereto in accordance with one or ³⁵ more aspects of the present disclosure.

FIGS. 51 and 52 illustrate one aspect of a distal sensor plug where FIG. 51 illustrates a cutaway view of the distal sensor plug and FIG. 52 further illustrates the magnetic field sensor and the processor operatively coupled to the flex board such that they are capable of communicating in accordance with one or more aspects of the present disclosure.

FIG. 53 illustrates an aspect of an end effector with a flex cable operable to provide power to sensors and electronics in the distal tip of the anvil portion in accordance with one or more aspects of the present disclosure.

FIG. 54 is a perspective view of an end effector of a surgical stapling instrument including a cartridge ⁵⁰ channel, a staple cartridge positioned in the cartridge channel, and an anvil in accordance with one or more aspects of the present disclosure.

FIG. 55 is a cross-sectional elevational view of the surgical stapling instrument of FIG. 134 illustrating a sled and a firing member in an unfired position in accordance with one or more aspects of the present disclosure.

FIG. 56 is a detail view depicting the sled of FIG. 55 in a partially advanced position and the firing member in its unfired position in accordance with one or more aspects of the present disclosure.

FIG. 57 illustrates one aspect of an end effector comprising a first sensor and a second sensor in accordance with one or more aspects of the present disclosure.

FIG. 58 is a logic diagram illustrating one aspect of a process for determining the thickness of a tissue section clamped between an anvil and a staple cartridge of an end effector in accordance with one or more aspects of the present disclosure.

FIG. 59 is a logic diagram illustrating one aspect of a process for determining the thickness of a tissue section clamped between the anvil and the staple cartridge of the end effector in accordance with one or more aspects of the present disclosure.

FIG. 60 illustrates one aspect of an end effector comprising a first sensor and a second sensor in accordance with one or more aspects of the present disclosure.

FIG. 61 illustrates one aspect of an end effector comprising a first sensor and a plurality of second sensors in accordance with one or more aspects of the present disclosure.

FIG. 62 illustrates one aspect of an end effector comprising a plurality of sensors in accordance with one or more aspects of the present disclosure.

FIG. 63 is a logic diagram illustrating one aspect of a process for determining one or more tissue properties based on a plurality of sensors in accordance with one or more aspects of the present disclosure. FIG. 64 illustrates one aspect of an end effector comprising a plurality of sensors coupled to a jaw member in accordance with one or more aspects of the present disclosure.

FIG. 65 illustrates one aspect of a staple cartridge comprising a plurality of sensors formed integrally therein in accordance with one or more aspects of the present disclosure.

FIG. 66 is a logic diagram illustrating one aspect of a process for determining one or more parameters of a tissue section clamped within an end effector in accordance with one or more aspects of the present disclosure.

FIG. 67 illustrates one aspect of an end effector comprising a sensor comprising a specific sampling rate to limit or eliminate false signals in accordance with one or more aspects of the present disclosure.

FIG. 68 is a logic diagram illustrating one aspect of a process for generating a thickness measurement for a tissue section located between an anvil and a staple cartridge of an end effector in accordance with one or more aspects of the present disclosure.

FIGS. 69A-69B illustrate one aspect of an end effector comprising a pressure sensor in accordance with one or more aspects of the present disclosure.

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FIG. 70 illustrates one aspect of an end effector comprising a second sensor located between a staple cartridge and a jaw member in accordance with one or more aspects of the present disclosure.

FIG. 71 is a logic diagram illustrating one aspect of a process for determining the thickness of a tissue section clamped in an end effector, according to FIGS. 69A-69B or FIG. 70 in accordance with one or more aspects of the present disclosure.

FIG. 72 illustrates one aspect of an end effector comprising a plurality of second sensors located between a staple cartridge and an elongated channel in accordance with one or more aspects of the present disclosure.

FIGS. 73A and 73B further illustrate the effect of a full versus partial bite of tissue in accordance with one or more aspects of the present disclosure.

FIG. 74 illustrates an aspect of an end effector that is configured to determine the location of a cutting member or knife in accordance with one or more ²⁰ aspects of the present disclosure.

FIG. 75 illustrates an example of the code strip in operation with red LEDs and an infrared LED in accordance with one or more aspects of the present disclosure.

FIG. 76 illustrates a partial perspective view of an end effector of a surgical instrument comprising a staple cartridge in accordance with one or more aspects of the present disclosure.

FIG. 77 illustrates an elevational view of a portion of ³⁰ the end effector of FIG. 76 in accordance with one or more aspects of the present disclosure.

FIG. 78 illustrates a logic diagram of a module of the surgical instrument of FIG. 76 in accordance with one or more aspects of the present disclosure.

FIG. 79 illustrates a partial view of a cutting edge, an optical sensor, and a light source of the surgical instrument of FIG. 76 in accordance with one or more aspects of the present disclosure.

FIG. 80 illustrates a partial view of a cutting edge, an optical sensor, and a light source of the surgical instrument of FIG. 76 in accordance with one or more aspects of the present disclosure.

FIG. 81 illustrates a partial view of a cutting edge, an optical sensor, and a light source of the surgical instrument of FIG. 76 in accordance with one or more aspects of the present disclosure.

FIG. 82 illustrates a partial view of a cutting edge, optical sensors, and light sources of the surgical instrument of FIG. 76 in accordance with one or more 50 aspects of the present disclosure.

FIG. 83 illustrates a partial view of a cutting edge, an optical sensor, and a light source of the surgical instrument of FIG. 76 in accordance with one or more aspects of the present disclosure.

FIG. 84 illustrates a perspective view of a staple cartridge including a sharpness testing member in accordance with one or more aspects of the present disclosure.

FIG. 85 illustrates a logic diagram of a module of a surgical instrument in accordance with one or more aspects of the present disclosure.

FIG. 86 illustrates a logic diagram of a module of a surgical instrument in accordance with one or more aspects of the present disclosure.

FIG. 87 illustrates a logic diagram outlining a method for evaluating sharpness of a cutting edge of a surgical instrument in accordance with one or more aspects of the present disclosure.

FIG. 88 illustrates a flow chart outlining a method for determining whether a cutting edge of a surgical instrument is sufficiently sharp to transect tissue captured by the surgical instrument in accordance with

one or more aspects of the present disclosure.

FIG. 89 illustrates a table showing predefined tissue thicknesses and corresponding predefined threshold forces in accordance with one or more aspects of the present disclosure.

FIG. 90 illustrates a logic diagram of a common controller for use with a plurality of motors of a surgical instrument in accordance with one or more aspects of the present disclosure.

FIG. 91 illustrates a partial elevational view of the handle of the surgical instrument with a removed outer casing in accordance with one or more aspects of the present disclosure.

FIG. 92 illustrates a partial elevational view of the surgical instrument with a removed outer casing in accordance with one or more aspects of the present disclosure.

FIG. 93A illustrates a side angle view of an end effector with the anvil in a closed position, illustrating one located on either side of the cartridge deck in accordance with one or more aspects of the present disclosure.

FIG. 93B illustrates a three-quarter angle view of the end effector with the anvil in an open position, and one LED located on either side of the cartridge deck in accordance with one or more aspects of the present disclosure.

FIG. 94A illustrates a side angle view of an end effector with the anvil in a closed position and a plurality of LEDs located on either side of the cartridge deck in accordance with one or more aspects of the present disclosure.

FIG. 94B illustrates a three-quarter angle view of the end effector with the anvil in an open position, and a plurality of LEDs located on either side of the cartridge deck in accordance with one or more aspects of the present disclosure.

FIG. 95A illustrates a side angle view of an end effector with the anvil in a closed position, and a plurality of LEDs from the proximal to the distal end of the staple cartridge, on either side of the cartridge deck in accordance with one or more aspects of the present disclosure.

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FIG. 95B illustrates a three-quarter angle view of the end effector with the anvil in an open position, illustrating a plurality of LEDs from the proximal to the distal end of the staple cartridge, and on either side of the cartridge deck in accordance with one or more aspects of the present disclosure.

FIG. 96 is a circuit diagram of an example power assembly of a surgical instrument in accordance with one or more aspects of the present disclosure.

FIG. 97 is a circuit diagram of an example power assembly of a surgical instrument in accordance with one or more aspects of the present disclosure.

FIG. 98 is a schematic block diagram of a control system of a surgical instrument in accordance with one or more aspects of the present disclosure.

FIG. 99 is a schematic block diagram of a control system of a surgical instrument in accordance with one or more aspects of the present disclosure.

FIG. 100 is a schematic diagram of an absolute positioning system comprising a controlled motor drive circuit arrangement comprising a sensor arrangement in accordance with one or more aspects of the present disclosure.

FIG. 101 is a detail perspective view of a sensor arrangement for an absolute positioning system in accordance with one or more aspects of the present disclosure.

FIG. 102 is an exploded perspective view of the sensor arrangement for an absolute positioning system showing a control circuit board assembly and the relative alignment of the elements of the sensor arrangement in accordance with one or more aspects of the present disclosure.

FIG. 103 is a schematic diagram of one aspect of a position sensor for an absolute positioning system ³⁵ comprising a magnetic rotary absolute positioning system in accordance with one or more aspects of the present disclosure.

FIG. 104 is a schematic illustrating a system for controlling the speed of a motor and/or the speed of a ⁴⁰ drivable member of a surgical instrument in accordance with one or more aspects of the present disclosure.

FIG. 105 is a schematic illustrating another system for controlling the speed of a motor and/or the speed of a drivable member of a surgical instrument in accordance with one or more aspects of the present disclosure.

FIG. 106 is a side view of one aspect of a handle assembly of a modular surgical instrument in accordance with one or more aspects of the present disclosure.

FIG. 107 is a block diagram of an aspect of a control system of a surgical instrument in accordance with one or more aspects of the present disclosure.

FIG. 108 is a perspective view of an interface of a modular surgical instrument in accordance with one or more aspects of the present disclosure.

FIG. 109 is a plan view of the interface shown in FIG. 108 of a modular surgical instrument in accordance with one or more aspects of the present disclosure. FIG. 110 is a first side elevational view of the interface shown in FIG. 108 of a modular surgical instrument in accordance with one or more aspects of the present disclosure.

FIG. 111 is a second side elevational view of the interface shown in FIG. 108 of a modular surgical instrument in accordance with one or more aspects of the present disclosure.

FIG. 112 is a side elevational view of another aspect of a handle assembly of a modular surgical instrument in accordance with one or more aspects of the present disclosure.

FIG. 113 is a side elevational view of another aspect of a handle assembly of a modular surgical instrument in accordance with one or more aspects of the present disclosure.

FIG. 114 is a flowchart of a method of operating a surgical instrument in accordance with one or more aspects of the present disclosure.

FIG. 115 is a flowchart of another method of operating a surgical instrument in accordance with one or more aspects of the present disclosure.

DETAILED DESCRIPTION

[0007] Applicant of the present application owns the following patent applications that were filed on even date herewith and which are each herein incorporated by reference in their respective entireties:

U.S. Patent Application Serial No. 15/130575, entitled STAPLE FORMATION DETECTION MECHA-NISMS. Attorney Docket No. END7774USNP/150513; U.S. Patent Application Serial No. 15/130582, entitled SURGICAL INSTRUMENT WITH DETECTION SENSORS. Attorney Docket No END7775USNP/150514; U.S. Patent Application Serial No. 15/130588, entitled SURGICAL INSTRUMENT WITH IMPROVED STOP/START CONTROL DURING A FIRING MO-TION, Attorney Docket No. END7776USNP/150515; U.S. Patent Application Serial No. 15/130595, entitled SURGICAL INSTRUMENT WITH ADJUSTA-BLE STOP/START CONTROL DURING A FIRING MOTION, Attorney Docket No. END7777USNP/150516; U.S. Patent Application Serial No. 15/130566, entitled SURGICAL INSTRUMENT WITH MULTIPLE PROGRAM RESPONSES DURING A FIRING MO-TION. Attorney Docket No. END7782USNP/150517; U.S. Patent Application Serial No. 15/130571, entitled SURGICAL INSTRUMENT WITH MULTIPLE

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PROGRAM RESPONSES DURING A FIRING MO-TION, Attorney Docket No. END7783USNP/150518; U.S. Patent Application Serial No. 15/130590, entitled SYSTEMS AND METHODS FOR CONTROL-LING A SURGICAL STAPLING AND CUTTING IN-STRUMENT, Attorney Docket No. END7785USNP/150520; and

U.S. Patent Application Serial No. 15/130596, entitled SYSTEMS AND METHODS FOR CONTROL-LING A SURGICAL STAPLING AND CUTTING IN-STRUMENT, Attorney Docket No. END7786USNP/150521.

[0008] The present disclosure provides an overall understanding of the principles of the structure, function, manufacture, and use of the devices and methods disclosed herein. One or more examples of these aspects are illustrated in the accompanying drawings. Those of ordinary skill in the art will understand that the devices and methods specifically described herein and illustrated in the accompanying are non-limiting examples. The features illustrated or described in connection with one example may be combined with the features of other examples. Such modifications and variations are intended to be included within the scope of the present disclosure.

[0009] Various example devices and methods are provided for performing laparoscopic and minimally invasive surgical procedures. However, the person of ordinary skill in the art will readily appreciate that the various methods and devices disclosed herein can be used in numerous surgical procedures and applications including, for example, in connection with open surgical procedures. As the present Detailed Description proceeds, those of ordinary skill in the art will further appreciate that the various instruments disclosed herein can be inserted into a body in any way, such as through a natural orifice, through an incision or puncture hole formed in tissue, etc. The working portions or end effector portions of the instruments can be inserted directly into a patient's body or can be inserted through an access device that has a working channel through which the end effector and elongated shaft of a surgical instrument can be advanced.

[0010] In one aspect, a motorized surgical stapling and cutting instrument provides a user actuatable control which overrides any limitation imposed by the instrument. In one aspect, the user control overrides any automatic operational wait or speed change. In another aspect, the user control provides pre-selection as part of display (LCD) setup to ignore all triggers. A secondary button may be pressed in combination with a regular control button to override or stop automatic modification of motions related to that control. Furthermore, the motorized surgical stapling and cutting instrument provides adaptive speed control, such as, for example, when a clinical algorithm has determined that an automatic (not prompted by the user) movement is necessary. In another as-

pect, a controller re-links the articulation mechanism to the firing mechanism after a transection. If a user has not pressed any related controls, the target speed during the movement is a first pre-defined value. In yet another aspect, when a user has not pressed any controls related to articulation, a low speed is selected. While the drive train is moving at the first target speed to the target position, if a user presses a related control, a second target speed is selected. For example, when the user presses

¹⁰ the articulate left button, a faster speed is selected such that the articulation mechanism is linked quicker, and the articulate left function is more readily available. The above example, when used on the motorized surgical stapling and cutting instrument described herein, pro-¹⁵ vides balance concerning the user, minimizing un-

prompted motor movement noise, and delaying the procedure, a second higher speed is initiated when the user selects a related articulation sweep control.

[0011] Before describing various aspects of a motor-20 ized stapling and cutting instrument (surgical instrument) as described in connection with FIGS. 106-115, the present disclosure first turns to FIGS. 1-105 for a general description of the mechanical and electrical platform upon which the present motorized surgical instrument may 25 be implemented and provides the background necessary to appreciate the underlying operation and functionality of the motorized surgical instrument. Accordingly, FIGS. 1-14 provide an example of a general description of the underlying mechanical platform upon which the present 30 motorized stapling and cutting instrument may be implemented. FIGS. 15-21 describe examples of the general underlying microcontroller, motor drive, and electrical interconnection platform upon which the present motorized surgical instrument may be implemented. FIGS. 22-34

- ³⁵ describe example end effector channel frames and measuring forces applied to tissue located between the anvil and the staple cartridge of the end effector. FIGS. 35-37 described example circuits for controlling the functionality of the present motorized surgical instrument.
- 40 FIGS. 38-95 describe example sensors and feedback systems to utilize the sensors outputs to implement the present motorized surgical instrument. FIGS. 97-97 describe example power assemblies for powering the present motorized surgical instrument. FIGS. 98-105 de-
- ⁴⁵ scribe example control systems for controlling motor speed and drivable members of the present surgical instrument includes sensors and feedback elements therefor. Upon familiarization with the underlying mechanical and electrical platform upon which the present motorized
- 50 surgical instrument may be implemented, the reader is directed to the description in connection with FIGS. 106-115 for a description of a motorized modular surgical stapling and cutting instrument with configurable operating mode.
- ⁵⁵ [0012] Accordingly, turning now to the figures, FIGS.
 1-6 depict a motor-driven surgical instrument 10 for cutting and fastening that may or may not be reused. In the illustrated examples, the surgical instrument 10 includes

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a housing 12 that comprises a handle assembly 14 that is configured to be grasped, manipulated and actuated by the clinician. The housing 12 is configured for operable attachment to an interchangeable shaft assembly 200 that has an end effector 300 operably coupled thereto that is configured to perform one or more surgical tasks or procedures. As the present Detailed Description proceeds, it will be understood that the various unique and novel arrangements of the various forms of interchangeable shaft assemblies disclosed herein also may be effectively employed in connection with robotically-controlled surgical systems. Thus, the term "housing" also may encompass a housing or similar portion of a robotic system that houses or otherwise operably supports at least one drive system that is configured to generate and apply at least one control motion which could be used to actuate the interchangeable shaft assemblies disclosed herein and their respective equivalents. The term "frame" may refer to a portion of a handheld surgical instrument. The term "frame" also may represent a portion of a robotically controlled surgical instrument and/or a portion of the robotic system that may be used to operably control a surgical instrument. For example, the interchangeable shaft assemblies disclosed herein may be employed with various robotic systems, instruments, components and methods disclosed in U.S. Patent No. 9,072,535, entitled SURGICAL STAPLING INSTRUMENTS WITH ROTAT-ABLE STAPLE DEPLOYMENT ARRANGEMENTS, which is incorporated by reference herein in its entirety. [0013] The housing 12 depicted in FIGS. 1-2 is shown in connection with an interchangeable shaft assembly 200 that includes an end effector 300 that comprises a surgical cutting and fastening device that is configured to operably support a surgical staple cartridge 304 therein. The housing 12 may be configured for use in connection with interchangeable shaft assemblies that include end effectors that are adapted to support different sizes and types of staple cartridges, have different shaft lengths, sizes, and types, etc. In addition, the housing 12 also may be effectively employed with a variety of other interchangeable shaft assemblies including those assemblies that are configured to apply other motions and forms of energy such as, for example, radio frequency (RF) energy, ultrasonic energy and/or motion to end effector arrangements adapted for use in connection with various surgical applications and procedures. Furthermore, the end effectors, shaft assemblies, handles, surgical instruments, and/or surgical instrument systems can utilize any suitable fastener, or fasteners, to fasten tissue. For instance, a fastener cartridge comprising a plurality of fasteners removably stored therein can be removably inserted into and/or attached to the end effector of a shaft assembly.

[0014] FIG. 1 illustrates the surgical instrument 10 with an interchangeable shaft assembly 200 operably coupled thereto. FIG. 2 illustrates attachment of the interchangeable shaft assembly 200 to the housing 12 or handle assembly 14. As shown in FIG. 4, the handle assembly 14 may comprise a pair of interconnectable handle housing segments 16 and 18 that may be interconnected by screws, snap features, adhesive, etc. In the illustrated arrangement, the handle housing segments 16, 18 cooperate to form a pistol grip portion 19 that can be gripped and manipulated by the clinician. As will be discussed in further detail below, the handle assembly 14 operably supports a plurality of drive systems therein that are configured to generate and apply various control motions to corresponding portions of the interchangeable shaft as-

sembly that is operably attached thereto.[0015] Referring now to FIG. 4, the handle assembly 14 may further include a frame 20 that operably supports a plurality of drive systems. For example, the frame 20

can operably support a "first" or closure drive system, generally designated as 30, which may be employed to apply closing and opening motions to the interchangeable shaft assembly 200 that is operably attached or coupled thereto. In at least one form, the closure drive system
 30 may include an actuator in the form of a closure trigger 32 that is pivotally supported by the frame 20. More spe-

cifically, as illustrated in FIG. 4, the closure trigger 32 is pivotally coupled to the handle assembly 14 by a pivot pin 33. Such arrangement enables the closure trigger 32 to be manipulated by a clinician such that when the cli-

²⁰ to be manipulated by a clinician such that when the clinician grips the pistol grip portion 19 of the handle assembly 14, the closure trigger 32 may be easily pivoted from a starting or "unactuated" position to an "actuated" position and more particularly to a fully compressed or
³⁰ fully actuated position. The closure trigger 32 may be biased into the unactuated position by spring or other biasing arrangement (not shown). In various forms, the closure drive system 30 further includes a closure linkage assembly 34 that is pivotally coupled to the closure trigger 32. As shown in FIG. 4, the closure linkage assembly 34 may include a first closure link 36 and a second closure linkage

link 38 that are pivotally coupled to the closure trigger 32 by a pin 35. The second closure link 38 also may be referred to herein as an "attachment member" and include a transverse attachment pin 37.

[0016] Still referring to FIG. 4, it can be observed that the first closure link 36 may have a an end or locking wall 39 thereon that is configured to cooperate with a closure release assembly 60 that is pivotally coupled to the frame

45 20. In at least one form, the closure release assembly 60 may comprise a closure release button assembly 62 that has a distally protruding locking pawl 64 formed thereon. The closure release button assembly 62 may be pivoted in a counterclockwise direction by a release spring (not

⁵⁰ shown). As the clinician depresses the closure trigger 32 from its unactuated position towards the pistol grip portion 19 of the handle assembly 14, the first closure link 36 pivots upward to a point wherein the locking pawl 64 drops into retaining engagement with the locking wall 39 on the first closure link 36 thereby preventing the closure trigger 32 from returning to the unactuated position. Thus, the closure release assembly 60 serves to lock the closure trigger 32 in the fully actuated position. When the

clinician desires to unlock the closure trigger 32 to permit it to be biased to the unactuated position, the clinician simply pivots the closure release button assembly 62 such that the locking pawl 64 is moved out of engagement with the locking wall 39 on the first closure link 36. When the locking pawl 64 has been moved out of engagement with the first closure link 36, the closure trigger 32 may pivot back to the unactuated position. Other closure trigger locking and release arrangements also may be employed.

[0017] Further to the above, FIGS. 10-11 illustrate the closure trigger 32 in its unactuated position which is associated with an open, or unclamped, configuration of the interchangeable shaft assembly 200 in which tissue can be positioned between the jaws of the interchangeable shaft assembly 200. FIG. 12 illustrates the closure trigger 32 in its actuated position which is associated with a closed, or clamped, configuration of the interchangeable shaft assembly 200 in which tissue is clamped between the jaws of the interchangeable shaft assembly 200. Upon comparing FIGS. 11 and 13, the reader will appreciate that, when the closure trigger 32 is moved from its unactuated position (FIG. 11) to its actuated position (FIG. 13), the closure release button assembly 62 is pivoted between a first position (FIG. 11) and a second position (FIG. 13). The rotation of the closure release button assembly 62 can be referred to as being an upward rotation; however, at least a portion of the closure release button assembly 62 is being rotated toward the circuit board 100. Referring to FIG. 4, the closure release button assembly 62 can include an arm 61 extending therefrom and a magnetic element 63, such as a permanent magnet, for example, mounted to the arm 61. When the closure release button assembly 62 is rotated from its first position to its second position, the magnetic element 63 can move toward the circuit board 100. The circuit board 100 can include at least one sensor configured to detect the movement of the magnetic element 63. In at least one aspect, a magnetic field sensor 65, for example, can be mounted to the bottom surface of the circuit board 100. The magnetic field sensor 65 can be configured to detect changes in a magnetic field surrounding the magnetic field sensor 65 caused by the movement of the magnetic element 63. The magnetic field sensor 65 can be in signal communication with a controller 1500, for example, which can determine whether the closure release button assembly 62 is in its first position, which is associated with the unactuated position of the closure trigger 32 and the open configuration of the end effector, its second position, which is associated with the actuated position of the closure trigger 32 and the closed configuration of the end effector, and/or any position between the first position and the second position.

[0018] As used throughout the present disclosure, a magnetic field sensor may be a Hall effect sensor, search coil, fluxgate, optically pumped, nuclear precession, SQUID, Hall-effect, anisotropic magnetoresistance, giant magnetoresistance, magnetic tunnel junctions, giant

magnetoimpedance, magnetostrictive/piezoelectric composites, magnetodiode, magnetotransistor, fiber optic, magnetooptic, and microelectromechanical systemsbased magnetic sensors, among others.

⁵ **[0019]** In at least one form, the handle assembly 14 and the frame 20 may operably support another drive system referred to herein as a firing drive system 80 that is configured to apply firing motions to corresponding portions of the interchangeable shaft assembly attached

¹⁰ thereto. The firing drive system may 80 also be referred to herein as a "second drive system". The firing drive system 80 may employ an electric motor 82, located in the pistol grip portion 19 of the handle assembly 14. In various forms, the electric motor 82 may be a DC brushed

¹⁵ driving motor having a maximum rotation of, approximately, 25,000 RPM, for example. In other arrangements, the motor may include a brushless motor, a cordless motor, a synchronous motor, a stepper motor, or any other suitable electric motor. The electric motor 82

²⁰ may be powered by a power source 90 that in one form may comprise a removable power pack 92. As shown in FIG. 4, for example, the removable power pack 92 may comprise a proximal housing portion 94 that is configured for attachment to a distal housing portion 96. The proxi-

mal housing portion 94 and the distal housing portion 96 are configured to operably support a plurality of batteries 98 therein. Batteries 98 may each comprise, for example, a Lithium Ion ("LI") or other suitable battery. The distal housing portion 96 is configured for removable operable
attachment to a control circuit board 100 which is also operably coupled to the electric motor 82. A number of batteries 98 may be connected in series may be used as the power source for the surgical instrument 10. In addition, the power source 90 may be replaceable and/or rechargeable.

[0020] As outlined above with respect to other various forms, the electric motor 82 can include a rotatable shaft (not shown) that operably interfaces with a gear reducer assembly 84 that is mounted in meshing engagement
with a with a set, or rack, of drive teeth 122 on a longitudinally movable drive member 120. In use, a voltage polarity provided by the power source 90 can operate the electric motor 82 in a clockwise direction wherein the voltage polarity applied to the electric motor by the battery
can be reversed in order to operate the electric motor 82

in a counter-clockwise direction. When the electric motor 82 is rotated in one direction, the longitudinally movable drive member 120 will be axially driven in the distal direction "DD". When the electric motor 82 is driven in the opposite rotary direction, the longitudinally movable drive member 120 will be axially driven in a proximal direction "PD". The handle assembly 14 can include a switch which can be configured to reverse the polarity applied to the electric motor 82 by the power source 90. As with the other forms described herein, the handle assembly 14 can also include a sensor that is configured to detect the position of the longitudinally movable drive member 120 and/or the direction in which the longitudinally movable

drive member 120 is being moved.

[0021] Actuation of the electric motor 82 can be controlled by a firing trigger 130 that is pivotally supported on the handle assembly 14. The firing trigger 130 may be pivoted between an unactuated position and an actuated position. The firing trigger 130 may be biased into the unactuated position by a spring 132 or other biasing arrangement such that when the clinician releases the firing trigger 130, it may be pivoted or otherwise returned to the unactuated position by the spring 132 or biasing arrangement. In at least one form, the firing trigger 130 can be positioned "outboard" of the closure trigger 32 as was discussed above. In at least one form, a firing trigger safety button 134 may be pivotally mounted to the closure trigger 32 by pin 35. The firing trigger safety button 134 may be positioned between the firing trigger 130 and the closure trigger 32 and have a pivot arm 136 protruding therefrom. See FIG. 4. When the closure trigger 32 is in the unactuated position, the firing trigger safety button 134 is contained in the handle assembly 14 where the clinician cannot readily access it and move it between a safety position preventing actuation of the firing trigger 130 and a firing position wherein the firing trigger 130 may be fired. As the clinician depresses the closure trigger 32, the firing trigger safety button 134 and the firing trigger 130 pivot down wherein they can then be manipulated by the clinician.

[0022] As discussed above, the handle assembly 14 can include a closure trigger 32 and a firing trigger 130. Referring to FIGS. 11-13, the firing trigger 130 can be pivotably mounted to the closure trigger 32. The closure trigger 32 can include an arm 31 extending therefrom and the firing trigger 130 can be pivotably mounted to the arm 31 about a pivot pin 33. When the closure trigger 32 is moved from its unactuated position (FIG.11) to its actuated position (FIG. 13), the firing trigger 130 can descend downwardly, as outlined above. After the firing trigger safety button 134 has been moved to its firing position, referring primarily to FIG. 18A, the firing trigger 130 can be depressed to operate the motor of the surgical instrument firing system. In various instances, the handle assembly 14 can include a tracking system, such as system 800, for example, configured to determine the position of the closure trigger 32 and/or the position of the firing trigger 130. With primary reference to FIGS. 11 and 13, the tracking system 800 can include a magnetic element, such as magnet 802, for example, which is mounted to an arm 801 extending from the firing trigger 130. The tracking system 800 can comprise one or more sensors, such as a first magnetic field sensor 803 and a second magnetic field sensor 804, for example, which can be configured to track the position of the magnet 802. [0023] Upon comparing FIGS. 11 and 13, the reader will appreciate that, when the closure trigger 32 is moved from its unactuated position to its actuated position, the magnet 802 can move between a first position adjacent the first magnetic field sensor 803 and a second position adjacent the second magnetic field sensor 804.

[0024] Upon comparing FIGS. 11 and 13, the reader will further appreciate that, when the firing trigger 130 is moved from an unfired position (FIG. 11) to a fired position (FIG. 13), the magnet 802 can move relative to the second magnetic field sensor 804. The first and second magnetic field sensors 803, 804 can track the movement of the magnet 802 and can be in signal communication with a controller on the circuit board 100. With data from the first magnetic field sensor 803 and/or the second

¹⁰ magnetic field sensor 804, the controller can determine the position of the magnet 802 along a predefined path and, based on that position, the controller can determine whether the closure trigger 32 is in its unactuated position, its actuated position, or a position therebetween.

Similarly, with data from the first magnetic field sensor 803 and/or the second magnetic field sensor 804, the controller can determine the position of the magnet 802 along a predefined path and, based on that position, the controller can determine whether the firing trigger 130 is
in its unfired position, its fully fired position, or a position

therebetween.
[0025] As indicated above, in at least one form, the longitudinally movable drive member 120 has a rack of drive teeth 122 formed thereon for meshing engagement
²⁵ with a corresponding drive gear 86 of the gear reducer assembly 84. At least one form also includes a manually-actuatable bailout assembly 140 that is configured to enable the clinician to manually retract the longitudinally movable drive member 120 should the electric motor 82
³⁰ become disabled. The bailout assembly 140 may include a lever or handle assembly 14 that is configured to be

- manually pivoted into ratcheting engagement with teeth 124 also provided in the longitudinally movable drive member 120. Thus, the clinician can manually retract the longitudinally movable drive member 120 by using the handle assembly 14 to ratchet the longitudinally movable
- drive member 120 in the proximal direction "PD". U.S. Patent No. 8,608,045, entitled POWERED SURGICAL CUTTING AND STAPLING APPARATUS WITH MANU-
- 40 ALLY RETRACTABLE FIRING SYSTEM discloses bailout arrangements and other components, arrangements and systems that also may be employed with the various instruments disclosed herein. U.S. Patent No. 8,608,045, is herein incorporated by reference in its entirety.

45 [0026] Turning now to FIG. 1, the interchangeable shaft assembly 200 includes an end effector 300 that comprises an elongated channel 302 that is configured to operably support a surgical staple cartridge 304 therein. The end effector 300 may further include an anvil 306 50 that is pivotally supported relative to the elongated channel 302. The interchangeable shaft assembly 200 may further include an articulation joint 270 and an articulation lock 350 (FIG. 7) which can be configured to releasably hold the end effector 300 in a desired position relative to 55 a shaft axis SA-SA. Details regarding the construction and operation of the end effector 300, the articulation joint 270 and the articulation lock 350 are set forth in U.S. Patent Application Publication No. 2014/0263541, enti-

tled ARTICULATABLE SURGICAL INSTRUMENT COMPRISING AN ARTICULATION LOCK, which is herein incorporated by reference in its entirety. As shown in FIG. 7, the interchangeable shaft assembly 200 can further include a proximal housing or nozzle 201 comprised of nozzle portions 202, 203. The interchangeable shaft assembly 200 can further include a closure tube 260 which can be utilized to close and/or open the anvil 306 of the end effector 300. Primarily referring now to FIG. 7, the interchangeable shaft assembly 200 can include a spine 210 which can be configured to fixably support a shaft frame 212 of the articulation lock 350. See FIG. 7. The spine 210 can be configured to, one, slidably support a firing member 220 therein and, two, slidably support the closure tube 260 which extends around the spine 210. The spine 210 can also be configured to slidably support an articulation driver 230. The articulation driver 230 has a distal end 231 that is configured to operably engage the articulation lock 350. The articulation lock 350 interfaces with an articulation frame 352 that is adapted to operably engage a drive pin (not shown) on the end effector frame (not shown). As indicated above, further details regarding the operation of the articulation lock 350 and the articulation frame may be found in U.S. Patent Application Publication No. 2014/0263541. In various circumstances, the spine 210 can comprise a proximal end 211 which is rotatably supported in a chassis 240. In one arrangement, for example, the proximal end 211 of the spine 210 has a thread 214 formed thereon for threaded attachment to a spine bearing 216 configured to be supported within the chassis 240. Such an arrangement facilitates rotatable attachment of the spine 210 to the chassis 240 such that the spine 210 may be selectively rotated about a shaft axis SA-SA relative to the chassis 240.

[0027] The interchangeable shaft assembly 200 includes a closure shuttle 250 that is slidably supported within the chassis 240 such that it may be axially moved relative thereto. As shown in FIG. 3, the closure shuttle 250 includes a pair of proximally-protruding hooks 252 that are configured for attachment to the transverse attachment pin 37 that is attached to the second closure link 38 as will be discussed in further detail below. A proximal end 261 of the closure tube 260 is coupled to the closure shuttle 250 for relative rotation thereto. For example, a U shaped connector 263 is inserted into an annular slot 262 in the proximal end 261 of the closure tube 260 and is retained within vertical slots 253 in the closure shuttle 250. Such an arrangement serves to attach the closure tube 260 to the closure shuttle 250 for axial travel therewith while enabling the closure tube 260 to rotate relative to the closure shuttle 250 about the shaft axis SA-SA. A closure spring 268 is journaled on the closure tube 260 and serves to bias the closure tube 260 in the proximal direction "PD" which can serve to pivot the closure trigger into the unactuated position when the shaft assembly is operably coupled to the handle assembly 14. [0028] In at least one form, the interchangeable shaft

assembly 200 may further include an articulation joint 270. Other interchangeable shaft assemblies, however, may not be capable of articulation. According to various forms, the double pivot closure sleeve assembly 271 includes an end effector closure sleeve assembly 272 having upper and lower distally projecting tangs 273, 274.

An end effector closure sleeve assembly 272 includes a horseshoe aperture 275 and a tab 276 for engaging an opening tab on the anvil 306 in the various manners de-

¹⁰ scribed in U.S. Patent Application Publication No. 2014/0263541. As described in further detail therein, the horseshoe aperture 275 and tab 276 engage a tab on the anvil when the anvil 306 is opened. An upper double pivot link 277 includes upwardly projecting distal and

¹⁵ proximal pivot pins that engage respectively an upper distal pin hole in the upper proximally projecting tang 273 and an upper proximal pin hole in an upper distally projecting tang 264 on the closure tube 260. A lower double pivot link 278 includes upwardly projecting distal and ²⁰ proximal pivot pins that engage respectively a lower distal pin hole in the lower proximally projecting tang 274 and a lower proximal pin hole in the lower distally projecting tang 265. See also FIG. 7.

[0029] In use, the closure tube 260 is translated distally
(direction "DD") to close the anvil 306, for example, in response to the actuation of the closure trigger 32. The anvil 306 is closed by distally translating the closure tube 260 and thus the end effector closure sleeve assembly 272, causing it to strike a proximal surface on the anvil 306 in the manner described in the aforementioned reference U.S. Patent Application Publication No.

2014/0263541. As was also described in detail in that reference, the anvil 306 is opened by proximally translating the closure tube 260 and the end effector closure
sleeve assembly 272, causing tab 276 and the horseshoe aperture 275 to contact and push against the anvil tab to

lift the anvil 306. In the anvil-open position, the closure tube 260 is moved to its proximal position.

[0030] As indicated above, the surgical instrument 10 may further include an articulation lock 350 of the types and construction described in further detail in U.S. Patent Application Publication No. 2014/0263541, which can be configured and operated to selectively lock the end effector 300 in position. Such arrangement enables the end

⁴⁵ effector 300 to be rotated, or articulated, relative to the closure tube 260 when the articulation lock 350 is in its unlocked state. In such an unlocked state, the end effector 300 can be positioned and pushed against soft tissue and/or bone, for example, surrounding the surgical site within the patient in order to cause the end effector 300.

^o within the patient in order to cause the end effector 300 to articulate relative to the closure tube 260. The end effector 300 also may be articulated relative to the closure tube 260 by an articulation driver 230.

[0031] As was also indicated above, the interchangeable shaft assembly 200 further includes a firing member 220 that is supported for axial travel within the spine 210. The firing member 220 includes an intermediate firing shaft 222 that is configured for attachment to a distal

cutting portion or knife bar 280. The firing member 220 also may be referred to herein as a "second shaft" and/or a "second shaft assembly". As shown in FIG. 7, the intermediate firing shaft 222 may include a longitudinal slot 223 in the distal end thereof which can be configured to receive a tab 284 on the proximal end 282 of the knife bar 280. The longitudinal slot 223 and the proximal end 282 can be sized and configured to permit relative movement therebetween and can comprise a slip joint 286. The slip joint 286 can permit the intermediate firing shaft 222 of the firing member 220 to be moved to articulate the end effector 300 without moving, or at least substantially moving, the knife bar 280. Once the end effector 300 has been suitably oriented, the intermediate firing shaft 222 can be advanced distally until a proximal sidewall of the longitudinal slot 223 comes into contact with the tab 284 in order to advance the knife bar 280 and fire the staple cartridge positioned within the channel 302. As can be further seen in FIG. 7, the spine 210 has an elongated opening or window 213 therein to facilitate assembly and insertion of the intermediate firing shaft 222 into the spine 210. Once the intermediate firing shaft 222 has been inserted therein, a top frame segment 215 may be engaged with the shaft frame 212 to enclose the intermediate firing shaft 222 and knife bar 280 therein. Further description of the operation of the firing member 220 may be found in U.S. Patent Application Publication No. 2014/0263541.

[0032] Further to the above, the interchangeable shaft assembly 200 can include a clutch assembly 400 which can be configured to selectively and releasably couple the articulation driver 230 to the firing member 220. In one form, the clutch assembly 400 includes a lock collar, or lock sleeve 402, positioned around the firing member 220 wherein the lock sleeve 402 can be rotated between an engaged position in which the lock sleeve 402 couples the articulation driver 360 to the firing member 220 and a disengaged position in which the articulation driver 360 is not operably coupled to the firing member 220. When lock sleeve 402 is in its engaged position, distal movement of the firing member 220 can move the articulation driver 360 distally and, correspondingly, proximal movement of the firing member 220 can move the articulation driver 230 proximally. When lock sleeve 402 is in its disengaged position, movement of the firing member 220 is not transmitted to the articulation driver 230 and, as a result, the firing member 220 can move independently of the articulation driver 230. In various circumstances, the articulation driver 230 can be held in position by the articulation lock 350 when the articulation driver 230 is not being moved in the proximal or distal directions by the firing member 220.

[0033] As shown in FIGS. 7-9, the interchangeable shaft assembly 200 further includes a switch drum 500 that is rotatably received on the closure tube 260. The switch drum 500 comprises a hollow shaft segment 502 that has a shaft boss 504 formed thereon for receive an outwardly protruding actuation pin 410 therein. In various

circumstances, the actuation pin 410 extends through a slot 267 into a longitudinal slot 408 provided in the lock sleeve 402 to facilitate axial movement of the lock sleeve 402 when it is engaged with the articulation driver 230. A rotary torsion spring 420 is configured to engage the shaft boss 504 on the switch drum 500 and a portion of

the nozzle portion 203 as shown in FIG. 8 to apply a biasing force to the switch drum 500. The switch drum 500 can further comprise at least partially circumferential openings 506 defined therein which, referring to FIGS. 5

¹⁰ openings 506 defined therein which, referring to FIGS. 5 and 6, can be configured to receive circumferential mounts 204, 205 extending from the nozzle portions 202, 203 and permit relative rotation, but not translation, between the switch drum 500 and the nozzle 201. As shown

¹⁵ in those Figures, the circumferential mounts 204, 205 also extend through openings 266 in the closure tube 260 to be seated in recesses located in the spine 210. However, rotation of the nozzle 201 to a point where the circumferential mounts 204, 205 reach the end of their

²⁰ respective partially circumferential openings 506 in the switch drum 500 will result in rotation of the switch drum 500 about the shaft axis SA-SA. Rotation of the switch drum 500 will ultimately result in the rotation of the actuation pin 410 and the lock sleeve 402 between its en-

²⁵ gaged and disengaged positions. Thus, in essence, the nozzle 201 may be employed to operably engage and disengage the articulation drive system with the firing drive system in the various manners described in further detail in U.S. Patent Application Publication No.
 30 2014/0263541.

[0034] As also illustrated in FIGS. 7-9, the interchangeable shaft assembly 200 can comprise a slip ring assembly 600 which can be configured to conduct electrical power to and/or from the end effector 300 and/or communicate signals to and/or from the end effector 300, for example. The slip ring assembly 600 can comprise a proximal connector flange 604 mounted to a chassis mounting flange 242 extending from the chassis 240 and a distal connector flange 601 positioned within a slot defined in the nozzle portions 202, 203. The proximal con-

nector flange 604 can comprise a first face and the distal connector flange 601 can comprise a second face which is positioned adjacent to and movable relative to the first face. The distal connector flange 601 can rotate relative

45 to the proximal connector flange 604 about the shaft axis SA-SA. The proximal connector flange 604 can comprise a plurality of concentric, or at least substantially concentric, conductors 602 defined in the first face thereof. A connector 607 can be mounted on the proximal side of 50 the distal connector flange 601 and may have a plurality of contacts (not shown) wherein each contact corresponds to and is in electrical contact with one of the conductors 602. Such an arrangement permits relative rotation between the proximal connector flange 604 and the 55 distal connector flange 601 while maintaining electrical contact therebetween. The proximal connector flange 604 can include an electrical connector 606 which can place the conductors 602 in signal communication with

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a shaft circuit board 610 mounted to the chassis 240, for example. In at least one instance, a wiring harness comprising a plurality of conductors can extend between the electrical connector 606 and the shaft circuit board 610. The electrical connector 606 may extend proximally through a connector opening 243 defined in the chassis mounting flange 242. U.S. Patent Application Publication No. 2014/0263551, entitled STAPLE CARTRIDGE TIS-SUE THICKNESS SENSOR SYSTEM, is incorporated herein by reference in its entirety. U.S. Patent Application Publication No. 2014/0263552, entitled STAPLE CAR-TRIDGE TISSUE THICKNESS SENSOR SYSTEM, is incorporated by reference in its entirety. Further details regarding slip ring assembly 600 may be found in U.S. Patent Application Publication No. 2014/0263541.

[0035] As discussed above, the interchangeable shaft assembly 200 can include a proximal portion which is fixably mounted to the handle assembly 14 and a distal portion which is rotatable about a longitudinal axis. The rotatable distal shaft portion can be rotated relative to the proximal portion about the slip ring assembly 600, as discussed above. The distal connector flange 601 of the slip ring assembly 600 can be positioned within the rotatable distal shaft portion. Moreover, further to the above, the switch drum 500 can also be positioned within the rotatable distal shaft portion. When the rotatable distal shaft portion is rotated, the distal connector flange 601 and the switch drum 500 can be rotated synchronously with one another. In addition, the switch drum 500 can be rotated between a first position and a second position relative to the distal connector flange 601. When the switch drum 500 is in its first position, the articulation drive system may be operably disengaged from the firing drive system and, thus, the operation of the firing drive system may not articulate the end effector 300 of the interchangeable shaft assembly 200. When the switch drum 500 is in its second position, the articulation drive system may be operably engaged with the firing drive system and, thus, the operation of the firing drive system may articulate the end effector 300 of the interchangeable shaft assembly 200. When the switch drum 500 is moved between its first position and its second position, the switch drum 500 is moved relative to distal connector flange 601. In various instances, the interchangeable shaft assembly 200 can comprise at least one sensor configured to detect the position of the switch drum 500. Turning now to FIGS 9, the distal connector flange 601 can comprise a magnetic field sensor 605, for example, and the switch drum 500 can comprise a magnetic element, such as permanent magnet 505, for example. The magnetic field sensor 605 can be configured to detect the position of the permanent magnet 505. When the switch drum 500 is rotated between its first position and its second position, the permanent magnet 505 can move relative to the magnetic field sensor 605. In various instances, magnetic field sensor 605 can detect changes in a magnetic field created when the permanent magnet 505 is moved. The magnetic field sensor 605 can be in signal communication with the shaft circuit board 610 and/or the circuit board 100 located in the handle, for example. Based on the signal from the magnetic field sensor 605, a controller on the shaft circuit board 610 and/or the circuit board 100 located in the handle can determine whether the articulation drive system is engaged with or disengaged from the firing drive system.

[0036] Referring again to FIG 3, the chassis 240 includes at least one, and preferably two, tapered attachment portions 244 formed thereon that are adapted to be

received within corresponding dovetail slots 702 formed within a distal attachment flange 700 of the frame 20. Each dovetail slot 702 may be tapered or, stated another way, be somewhat V-shaped to seatingly receive the ta-

pered attachment portions 244 therein. As can be further seen in FIG. 3, a shaft attachment lug 226 is formed on the proximal end of the intermediate firing shaft 222. As will be discussed in further detail below, when the interchangeable shaft assembly 200 is coupled to the handle
assembly 14, the shaft attachment lug 226 is received in a firing shaft attachment cradle 126 formed in the distal end 125 of the longitudinally movable drive member 120

as shown in FIGS. 3 and 6, for example. [0037] Various shaft assemblies employ a latch sys-25 tem 710 for removably coupling the interchangeable shaft assembly 200 to the housing 12 and more specifically to the frame 20. The proximally protruding lock lugs 714 each have a pivot lock lugs 716 formed thereon that are adapted to be received in corresponding holes 245 30 formed in the chassis 240. Such arrangement facilitates pivotal attachment of the lock yoke 712 to the chassis 240. The lock yoke 712 may include two proximally protruding lock lugs 714 that are configured for releasable engagement with corresponding lock detents or grooves 35 704 in the distal attachment flange 700 of the frame 20. See FIG. 3. In various forms, the lock yoke 712 is biased in the proximal direction by spring or biasing member

(not shown). Actuation of the lock yoke 712 may be accomplished by a latch button 722 that is slidably mounted
on a latch actuator assembly 720 that is mounted to the chassis 240. The latch button 722 may be biased in a

proximal direction relative to the lock yoke 712. As will be discussed in further detail below, the lock yoke 712 may be moved to an unlocked position by biasing the ⁴⁵ latch button the in distal direction which also causes the

lock yoke 712 to pivot out of retaining engagement with the distal attachment flange 700 of the frame 20. When the lock yoke 712 is in "retaining engagement" with the distal attachment flange 700 of the frame 20, the pivot
lock lugs 716 are retainingly seated within the corresponding lock detents or grooves 704 in the distal attachment flange 700.

[0038] When employing an interchangeable shaft assembly that includes an end effector of the type described
 ⁵⁵ herein that is adapted to cut and fasten tissue, as well as other types of end effectors, it may be desirable to prevent inadvertent detachment of the interchangeable shaft assembly from the housing during actuation of the

end effector. For example, in use the clinician may actuate the closure trigger 32 to grasp and manipulate the target tissue into a desired position. Once the target tissue is positioned within the end effector 300 in a desired orientation, the clinician may then fully actuate the closure trigger 32 to close the anvil 306 and clamp the target tissue in position for cutting and stapling. In that instance, the first drive system 30 has been fully actuated. After the target tissue has been clamped in the end effector 300, it may be desirable to prevent the inadvertent detachment of the interchangeable shaft assembly 200 from the housing 12. One form of the latch system 710 is configured to prevent such inadvertent detachment.

[0039] The lock voke 712 includes at least one, and preferably two, lock hooks 718 that are adapted to contact lock lugs 256 that are formed on the closure shuttle 250. Referring to FIGS. 10 and 11, when the closure shuttle 250 is in an unactuated position (i.e., the first closure drive system 30 is unactuated and the anvil 306 is open), the lock yoke 712 may be pivoted in a distal direction to unlock the interchangeable shaft assembly 200 from the housing 12. When in that position, the lock hooks 718 do not contact the lock lugs 256 on the closure shuttle 250. However, when the closure shuttle 250 is moved to an actuated position (i.e., the first closure drive system 30 is actuated and the anvil 306 is in the closed position), the lock yoke 712 is prevented from being pivoted to an unlocked position. See FIGS. 12 and 13. Stated another way, if the clinician were to attempt to pivot the lock yoke 712 to an unlocked position or, for example, the lock yoke 712 was in advertently bumped or contacted in a manner that might otherwise cause it to pivot distally, the lock hooks 718 on the lock yoke 712 will contact the lock lugs 256 on the closure shuttle 250 and prevent movement of the lock yoke 712 to an unlocked position.

[0040] Attachment of the interchangeable shaft assembly 200 to the handle assembly 14 will now be described with reference to FIG. 3. To commence the coupling process, the clinician may position the chassis 240 of the interchangeable shaft assembly 200 above or adjacent to the distal attachment flange 700 of the frame 20 such that the tapered attachment portions 244 formed on the chassis 240 are aligned with the dovetail slots 702 in the frame 20. The clinician may then move the interchangeable shaft assembly 200 along an installation axis IA that is perpendicular to the shaft axis SA-SA to seat the tapered attachment portions 244 in "operable engagement" with the corresponding dovetail receiving slots 702. In doing so, the shaft attachment lug 226 on the intermediate firing shaft 222 will also be seated in the firing shaft attachment cradle 126 in the longitudinally movable drive member 120 and the portions of the transverse attachment pin 37 on the second closure link 38 will be seated in the corresponding proximally-protruding hooks 252 in the closure shuttle 250. As used herein, the term "operable engagement" in the context of two components means that the two components are sufficiently engaged with each other so that upon application of an actuation motion thereto, the components may carry out their intended action, function and/or procedure.

- **[0041]** As discussed above, at least five systems of the interchangeable shaft assembly 200 can be operably coupled with at least five corresponding systems of the handle assembly 14. A first system can comprise a frame system which couples and/or aligns the frame or spine of the interchangeable shaft assembly 200 with the frame 20 of the handle assembly 14. Another system can com-
- ¹⁰ prise a closure drive system 30 which can operably connect the closure trigger 32 of the handle assembly 14 and the closure tube 260 and the anvil 306 of the interchangeable shaft assembly 200. As outlined above, the closure shuttle 250 of the interchangeable shaft assem-

¹⁵ bly 200 can be engaged with the transverse attachment pin 37 on the second closure link 38. Another system can comprise the firing drive system 80 which can operably connect the firing trigger 130 of the handle assembly 14 with the intermediate firing shaft 222 of the interchange-20 able shaft assembly 200.

[0042] As outlined above, the shaft attachment lug 226 can be operably connected with the firing shaft attachment cradle 126 of the longitudinally movable drive member 120. Another system can comprise an electrical sys-

tem which can signal to a controller in the handle assembly 14, such as controller, for example, that a shaft assembly, such as the interchangeable shaft assembly 200, for example, has been operably engaged with the handle assembly 14 and/or, two, conduct power and/or communication signals between the interchangeable shaft assembly 200 and the handle assembly 14. For instance, the interchangeable shaft assembly 200 can include an electrical connector 1410 that is operably mounted to the shaft circuit board 610. The electrical connector 1410

located on the shaft is configured for mating engagement with an electrical connector 1400 on the circuit board 100 located in the handle. Further details regaining the circuitry and control systems may be found in U.S. Patent Application Publication No. 2014/0263541. The fifth system may consist of the latching system for releasably

locking the interchangeable shaft assembly 200 to the handle assembly 14. [0043] Referring to FIG. 14, a non-limiting form of the

end effector 300 is illustrated. As described above, the
end effector 300 may include the anvil 306 and the surgical staple cartridge 304. In this non-limiting example, the anvil 306 is coupled to an elongated channel 198. For example, apertures 199 can be defined in the elongated channel 198 which can receive pins 152 extending
from the anvil 306 and allow the anvil 306 to pivot from an open position to a closed position relative to the elongated channel 198 and surgical staple cartridge 304. In addition, FIG. 14 shows a firing bar 172, configured to longitudinally translate into the end effector 300. The fir-

⁵⁵ ing bar 172 may be constructed from one solid section, or in various examples, may include a laminate material comprising, for example, a stack of steel plates. A distally projecting end of the firing bar 172 can be attached to an

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E-beam 178 that can, among other things, assist in spacing the anvil 306 from a surgical staple cartridge 304 positioned in the elongated channel 198 when the anvil 306 is in a closed position. The E-beam 178 can also include a sharpened cutting edge 182 which can be used to sever tissue as the E-beam 178 is advanced distally by the firing bar 172. In operation, the E-beam 178 can also actuate, or fire, the surgical staple cartridge 304. The surgical staple cartridge 304 can include a molded cartridge body 194 that holds a plurality of staples 191 resting upon staple drivers 192 within respective upwardly open staple cavities 195. A wedge sled 190 is driven distally by the E-beam 178, sliding upon a cartridge tray 196 that holds together the various components of the surgical staple cartridge 304. The wedge sled 190 upwardly cams the staple drivers 192 to force out the staples 191 into deforming contact with the anvil 306 while a cutting edge 182 of the E-beam 178 severs clamped tissue. [0044] Further to the above, the E-beam 178 can include upper pins 180 which engage the anvil 306 during firing. The E-beam 178 can further include middle pins 184 and a bottom foot 186 which can engage various portions of the cartridge body 194, cartridge tray 196 and elongated channel 198. When a surgical staple cartridge 304 is positioned within the elongated channel 198, a slot 193 defined in the cartridge body 194 can be aligned with a longitudinal slot 197 defined in the cartridge tray 196 and a slot 189 defined in the elongated channel 198. In use, the E-beam 178 can slide through the aligned longitudinal slots 193, 197, and 189 wherein, as indicated in FIG. 14, the bottom foot 186 of the E-beam 178 can engage a groove running along the bottom surface of elongated channel 198 along the length of slot 189, the middle pins 184 can engage the top surfaces of cartridge tray 196 along the length of longitudinal slot 197, and the upper pins 180 can engage the anvil 306. In such circumstances, the E-beam 178 can space, or limit the relative movement between, the anvil 306 and the surgical staple cartridge 304 as the firing bar 172 is moved distally to fire the staples from the surgical staple cartridge 304 and/or incise the tissue captured between the anvil 306 and the surgical staple cartridge 304. Thereafter, the firing bar 172 and the E-beam 178 can be retracted proximally allowing the anvil 306 to be opened to release the two stapled and severed tissue portions (not shown). [0045] Having described a surgical instrument 10 (FIGS. 1-14) in general terms, the description now turns to a detailed description of various electrical/electronic components of the surgical instrument 10. Referring again to FIGS. 2 and 3, the handle assembly 14 can include an electrical connector 1400 comprising a plurality of electrical contacts. Turning now to FIG. 15, the electrical connector 1400 can comprise a first electrical contact 1401a, a second electrical contact 1401b, a third electrical contact 1401c, a fourth electrical contact 1401d, a fifth electrical contact 1401e, and a sixth electrical contact 1401f, for example. While the illustrated example utilizes six contacts, other examples are envi-

sioned which may utilize more than six contacts or less than six contacts.

- [0046] As illustrated in FIG. 15, the first electrical contact 1401a can be in electrical communication with a transistor 1408, electrical contacts 1401b-1401e can be in electrical communication with a controller 1500, and the sixth electrical contact 1401f can be in electrical communication with a ground. In certain circumstances, one or more of the electrical contacts 1401b-1401e may be in
 10 electrical communication with one or more output chan
 - nels of the controller 1500 and can be energized, or have a voltage potential applied thereto, when the handle 1042 is in a powered state. In some circumstances, one or more of the electrical contacts 1401b-1401e may be in

¹⁵ electrical communication with one or more input channels of the controller 1500 and, when the handle assembly 14 is in a powered state, the controller 1500 can be configured to detect when a voltage potential is applied to such electrical contacts. When a shaft assembly, such as the

- ²⁰ interchangeable shaft assembly 200, for example, is assembled to the handle assembly 14, the electrical contacts 1401a-1401f may not communicate with each other. When a shaft assembly is not assembled to the handle assembly 14, however, the electrical contacts 1401a-
- 25 1401f of the electrical connector 1400 may be exposed and, in some circumstances, one or more of the electrical contacts 1401a-1401f may be accidentally placed in electrical communication with each other. Such circumstances can arise when one or more of the electrical contacts 30 1401a-1401f come into contact with an electrically conductive material, for example. When this occurs, the controller 1500 can receive an erroneous input and/or the interchangeable shaft assembly 200 can receive an erroneous output, for example. To address this issue, in 35 various circumstances, the handle assembly 14 may be unpowered when a shaft assembly, such as the interchangeable shaft assembly 200, for example, is not attached to the handle assembly 14.
- [0047] In other circumstances, the handle 1042 can be powered when a shaft assembly, such as the interchangeable shaft assembly 200, for example, is not attached thereto. In such circumstances, the controller 1500 can be configured to ignore inputs, or voltage potentials, applied to the contacts in electrical communica-
- tion with the controller 1500, i.e., electrical contacts 1401b-1401e, for example, until a shaft assembly is attached to the handle assembly 14. Even though the controller 1500 may be supplied with power to operate other functionalities of the handle assembly 14 in such circumstances, the handle assembly 14 may be in a powered-down state. In a way, the electrical connector 1400 may be in a powered-down state as voltage potentials applied
- to the electrical contacts 1401b-1401e may not affect the operation of the handle assembly 14. The reader will appreciate that, even though electrical contacts 1401b-1401e may be in a powered-down state, the electrical contacts 1401a and 1401f, which are not in electrical communication with the controller 1500, may or may not

be in a powered-down state. For instance, sixth electrical contact 1401f may remain in electrical communication with a ground regardless of whether the handle assembly 14 is in a powered-up or a powered-down state.

[0048] Furthermore, the transistor 1408, and/or any other suitable arrangement of transistors, such as transistor 1412, for example, and/or switches may be configured to control the supply of power from a power source 1404, such as a battery, within the handle assembly 14, for example, to the first electrical contact 1401a regardless of whether the handle assembly 14 is in a poweredup or a powered-down state. In various circumstances, the interchangeable shaft assembly 200, for example, can be configured to change the state of the transistor 1408 when the interchangeable shaft assembly 200 is engaged with the handle assembly 14. In certain circumstances, further to the below, a magnetic field sensor 1402 can be configured to switch the state of transistor 1412 which, as a result, can switch the state of transistor 1408 and ultimately supply power from power source 1404 to first electrical contact 1401a. In this way, both the power circuits and the signal circuits to the electrical connector 1400 can be powered down when a shaft assembly is not installed to the handle assembly 14 and powered up when a shaft assembly is installed to the handle assembly 14.

[0049] In various circumstances, referring again to FIG. 15, the handle assembly 14 can include the magnetic field sensor 1402, for example, which can be configured to detect a detectable element, such as a magnetic element 1407 (FIG. 3), for example, on a shaft assembly, such as the interchangeable shaft assembly 200, for example, when the shaft assembly is coupled to the handle assembly 14. The magnetic field sensor 1402 can be powered by a power source 1406, such as a battery, for example, which can, in effect, amplify the detection signal of the magnetic field sensor 1402 and communicate with an input channel of the controller 1500 via the circuit illustrated in FIG. 15. Once the controller 1500 has a received an input indicating that a shaft assembly has been at least partially coupled to the handle assembly 14, and that, as a result, the electrical contacts 1401a-1401f are no longer exposed, the controller 1500 can enter into its normal, or powered-up, operating state. In such an operating state, the controller 1500 will evaluate the signals transmitted to one or more of the electrical contacts 1401b-1401e from the shaft assembly and/or transmit signals to the shaft assembly through one or more of the electrical contacts 1401b-1401e in normal use thereof. In various circumstances, the interchangeable shaft assembly 200 may have to be fully seated before the magnetic field sensor 1402 can detect the magnetic element 1407. While a magnetic field sensor 1402 can be utilized to detect the presence of the interchangeable shaft assembly 200, any suitable system of sensors and/or switches can be utilized to detect whether a shaft assembly has been assembled to the handle assembly 14, for example. In this way, further to the above,

both the power circuits and the signal circuits to the electrical connector 1400 can be powered down when a shaft assembly is not installed to the handle assembly 14 and powered up when a shaft assembly is installed to the handle assembly 14.

[0050] In various examples, as may be used throughout the present disclosure, any suitable magnetic field sensor may be employed to detect whether a shaft assembly has been assembled to the handle assembly 14,

¹⁰ for example. For example, the technologies used for magnetic field sensing include Hall effect sensor, search coil, fluxgate, optically pumped, nuclear precession, SQUID (superconducting quantum interference device a very sensitive magnetometer used to measure ex-

¹⁵ tremely subtle magnetic fields, based on superconducting loops containing Josephson junctions), Hall-effect, anisotropic magnetoresistance, giant magnetoresistance, magnetic tunnel junctions, giant magnetoimpedance, magnetostrictive/piezoelectric composites, mag-

²⁰ netodiode, magnetotransistor, fiber optic, magnetooptic, and microelectromechanical systems-based magnetic sensors, among others.

[0051] Referring to FIG. 15, the controller 1500 may generally comprise a processor ("microprocessor") and
one or more memory units operationally coupled to the processor. By executing instruction code stored in the memory, the processor may control various components of the surgical instrument, such as the motor, various drive systems, and/or a user display, for example. The
controller 1500 may be implemented using integrated and/or discrete hardware elements, software elements, and/or a combination of both. Examples of integrated hardware elements may include processors, microprocessors, controllers, controllers, integrated circuits, appli-

 ³⁵ cation specific integrated circuits (ASIC), programmable logic devices (PLD), digital signal processors (DSP), field programmable gate arrays (FPGA), logic gates, registers, semiconductor devices, chips, microchips, chip sets, controllers, system-on-chip (SoC), and/or system ⁴⁰ in-package (SIP). Examples of discrete hardware ele-

ments may include circuits and/or circuit elements such as logic gates, field effect transistors, bipolar transistors, resistors, capacitors, inductors, and/or relays. In certain instances, the controller 1500 may include a hybrid circuit for comprising discrete and integrated circuit elements or

comprising discrete and integrated circuit elements or components on one or more substrates, for example. [0052] Referring to FIG. 15, the controller 1500 may be an LM4F230H5QR, available from Texas Instruments, for example. In certain instances, the Texas In-50 struments LM4F230H5QR is an ARM Cortex-M4F Processor Core comprising on-chip memory of 256 KB singlecycle flash memory, or other non-volatile memory, up to 40 MHz, a prefetch buffer to improve performance above 40 MHz, a 32 KB single-cycle serial random access mem-55 ory (SRAM), internal read-only memory (ROM) loaded with StellarisWare® software, 2KB electrically erasable programmable read-only memory (EEPROM), one or more pulse width modulation (PWM) modules, one or

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[0053] As discussed above, the handle assembly 14 and/or the interchangeable shaft assembly 200 can include systems and configurations configured to prevent, or at least reduce the possibility of, the contacts of the electrical connector 1400 located on the handle and/or the contacts of the electrical connector 1410 located on the shaft from becoming shorted out when the interchangeable shaft assembly 200 is not assembled, or completely assembled, to the handle assembly 14. Referring to FIG. 3, the electrical connector 1400 located on the handle can be at least partially recessed within a cavity 1409 defined in the frame 20. The six electrical contacts 1401a-1401f of the electrical connector 1400 can be completely recessed within the cavity 1409. Such arrangements can reduce the possibility of an object accidentally contacting one or more of the electrical contacts 1401a-1401f. Similarly, the electrical connector 1410 located on the shaft can be positioned within a recess defined in the chassis 240 which can reduce the possibility of an object accidentally contacting one or more of the electrical contacts 1411a-1411f of the electrical connector 1410 located on the shaft. With regard to the particular example depicted in FIG. 3, the electrical contacts 1411a-1411f located on the shaft can comprise male contacts. In at least one example, each of the electrical contacts 1411 a-1411f located in the shaft can comprise a flexible projection extending therefrom which can be configured to engage an electrical contact 1401a-1401f located on the handle, for example. The electrical contacts 1401a-1401f located on the handle can comprise female contacts. In at least one example, each electrical contact 1401a-1401f located on the handle can comprise a flat surface, for example, against which the male electrical contacts 1401a-1401f located on the shaft can wipe, or slide, against and maintain an electrically conductive interface therebetween. In various instances, the direction in which the interchangeable shaft assembly 200 is assembled to the handle assembly 14 can be parallel to, or at least substantially parallel to, the electrical contacts 1401a-1401f located on the handle such that the electrical contacts 1411a-1411f located on the shaft slide against the electrical contacts 1401a-1401f located on the handle when the interchangeable shaft assembly 200 is assembled to the handle assembly 14. In various alternative examples, the electrical contacts 1401a-1401f located in the handle can comprise male contacts and the electrical contacts 1411a-1411f located on the shaft can comprise female contacts. In certain alternative examples, the electrical contacts 1401a-1401f located on the handle and the electrical contacts 1411a-1411f located on the shaft can comprise any suitable arrangement of contacts.

[0054] In various instances, the handle assembly 14 can comprise a connector guard configured to at least partially cover the electrical connector 1400 located on the handle and/or a connector guard configured to at least partially cover the electrical connector 1410 located on the shaft. A connector guard can prevent, or at least reduce the possibility of, an object accidentally touching the contacts of an electrical connector when the shaft assembly is not assembled to, or only partially assembled to, the handle. A connector guard can be movable. For

instance, the connector guard can be moved between a guarded position in which it at least partially guards a connector and an unguarded position in which it does not guard, or at least guards less of, the connector. In at

least one example, a connector guard can be displaced as the shaft assembly is being assembled to the handle. For instance, if the handle comprises a handle connector guard, the shaft assembly can contact and displace the

²⁰ handle connector guard as the shaft assembly is being assembled to the handle. Similarly, if the shaft assembly comprises a shaft connector guard, the handle can contact and displace the shaft connector guard as the shaft assembly is being assembled to the handle. In various

25 instances, a connector guard can comprise a door, for example. In at least one instance, the door can comprise a beveled surface which, when contacted by the handle or shaft, can facilitate the displacement of the door in a certain direction. In various instances, the connector 30 guard can be translated and/or rotated, for example. In certain instances, a connector guard can comprise at least one film which covers the contacts of an electrical connector. When the shaft assembly is assembled to the handle, the film can become ruptured. In at least one 35 instance, the male contacts of a connector can penetrate the film before engaging the corresponding contacts positioned underneath the film.

[0055] As described above, the surgical instrument can include a system which can selectively power-up, or
 activate, the contacts of an electrical connector, such as the electrical connector 1400, for example. In various instances, the contacts can be transitioned between an unactivated condition and an activated condition. In certain instances, the contacts can be transitioned between

⁴⁵ a monitored condition, a deactivated condition, and an activated condition. For instance, the controller 1500, for example, can monitor the electrical contacts 1401a-1401f when a shaft assembly has not been assembled to the handle assembly 14 to determine whether one or
⁵⁰ more of the electrical contacts 1401a-1401f may have been shorted. The controller 1500 can be configured to apply a low voltage potential to each of the electrical contacts 1401a-1401f and assess whether only a minimal resistance is present at each of the contacts. Such an
⁵⁵ operating state can comprise the monitored condition. In

the event that the resistance detected at a contact is high, or above a threshold resistance, the controller 1500 can deactivate that contact, more than one contact, or, alter-

natively, all of the contacts. Such an operating state can comprise the deactivated condition. If a shaft assembly is assembled to the handle assembly 14 and it is detected by the controller 1500, as discussed above, the controller 1500 can increase the voltage potential to the electrical contacts 1401a-1401f. Such an operating state can comprise the activated condition.

[0056] The various shaft assemblies disclosed herein may employ sensors and various other components that require electrical communication with the controller in the housing. These shaft assemblies generally are configured to be able to rotate relative to the housing necessitating a connection that facilitates such electrical communication between two or more components that may rotate relative to each other. When employing end effectors of the types disclosed herein, the connector arrangements must be relatively robust in nature while also being somewhat compact to fit into the shaft assembly connector portion.

[0057] Turning now to FIGS. 16A and 16B, where one example of a segmented circuit 2000 comprising a plurality of circuit segments 2002a-2002g is illustrated. The segmented circuit 2000 comprising the plurality of circuit segments 2002a-2002g is configured to control a powered surgical instrument, such as, for example, the surgical instrument 10 illustrated in FIGS. 1-13, without limitation. The plurality of circuit segments 2002a-2002g is configured to control one or more operations of the powered surgical instrument 10. A safety processor segment 2002a (Segment 1) comprises a safety processor 2004. A primary processor segment 2002b (Segment 2) comprises a primary processor 2006. The safety processor 2004 and/or the primary processor 2006 are configured to interact with one or more additional circuit segments 2002c-2002g to control operation of the powered surgical instrument 10. The primary processor 2006 comprises a plurality of inputs coupled to, for example, one or more circuit segments 2002c-2002g, a battery 2008, and/or a plurality of switches 2058a-2070. The segmented circuit 2000 may be implemented by any suitable circuit, such as, for example, a printed circuit board assembly (PCBA) within the powered surgical instrument 10. It should be understood that the term processor as used herein includes any microprocessor, processors, controller, controllers, or other basic computing device that incorporates the functions of a computer's central processing unit (CPU) on an integrated circuit or at most a few integrated circuits. The processor is a multipurpose, programmable device that accepts digital data as input, processes it according to instructions stored in its memory, and provides results as output. It is an example of sequential digital logic, as it has internal memory. Processors operate on numbers and symbols represented in the binary numeral system.

[0058] In one aspect, the primary processor 2006 may be any single core or multicore processor such as those known under the trade name ARM Cortex by Texas Instruments. In one example, the safety processor 2004

may be a safety controller platform comprising two controller-based families such as TMS570 and RM4x known under the trade name Hercules ARM Cortex R4, also by Texas Instruments. Nevertheless, other suitable substitutes for controllers and safety processor may be em-

ployed, without limitation. In one example, the safety processor 2004 may be configured specifically for IEC 61508 and ISO 26262 safety critical applications, among others, to provide advanced integrated safety features

¹⁰ while delivering scalable performance, connectivity, and memory options. In certain instances, the primary processor 2006 may be a single core or multicore controller LM4F230H5QR as described in connection with FIGS. 14-17B.

¹⁵ [0059] In one aspect, the segmented circuit 2000 comprises an acceleration segment 2002c (Segment 3). The acceleration segment 2002c comprises an accelerometer 2022. The accelerometer 2022 is configured to detect movement or acceleration of the powered surgical instru-

²⁰ ment 10. In some examples, input from the accelerometer 2022 is used, for example, to transition to and from a sleep mode, identify an orientation of the powered surgical instrument, and/or identify when the surgical instrument has been dropped. In some examples, the accel-²⁵ eration segment 2002c is coupled to the safety processor

2004 and/or the primary processor 2006. **[0060]** In one aspect, the segmented circuit 2000 comprises a display segment 2002d (Segment 4). The display segment 2002d comprises a display connector 2024 cou-

pled to the primary processor 2006. The display connector 2024 couples the primary processor 2006 to a display 2028 through one or more integrated circuit drivers of the display 2026. The integrated circuit drivers of the display 2026 may be integrated with the display 2028 and/or may

³⁵ be located separately from the display 2028. The display 2028 may comprise any suitable display, such as, for example, an organic light-emitting diode (OLED) display, a liquid-crystal display (LCD), and/or any other suitable display. In some examples, the display segment 2002d
 ⁴⁰ is coupled to the safety processor 2004.

[0061] In some aspects, the segmented circuit 2000 comprises a shaft segment 2002e (Segment 5). The shaft segment 2002e comprises one or more controls for an interchangeable shaft assembly 200 (FIG. 1) coupled to the surgical instrument 10 and/or one or more controls for an end effector 300 coupled to the interchangeable

shaft assembly 200 (FIG. 1). The shaft segment 2002e comprises a shaft connector 2030 configured to couple the primary processor 2006 to a shaft PCBA 2031. The
shaft PCBA 2031 comprises a first articulation switch 2036, a second articulation switch 2032, and a shaft PCBA EEPROM 2034. In some examples, the shaft PCBA EEPROM 2034 comprises one or more parameters, routines, and/or programs specific to the interchangeable shaft assembly 200 and/or the shaft PCBA 2031. The shaft PCBA 2031 may be coupled to the interchangeable shaft assembly 200 and/or integral with the surgical instrument 10. In some examples, the shaft

segment 2002e comprises a second shaft EEPROM 2038. The second shaft EEPROM 2038 comprises a plurality of algorithms, routines, parameters, and/or other data corresponding to one or more shaft assemblies 200 and/or end effectors 300 which may be interfaced with the powered surgical instrument 10.

[0062] In some aspects, the segmented circuit 2000 comprises a position encoder segment 2002f (Segment 6). The position encoder segment 2002f comprises one or more magnetic angle rotary position encoders 2040a-2040b. The one or more magnetic angle rotary position encoders 2040a-2040b are configured to identify the rotational position of a motor 2048, an interchangeable shaft assembly 200 (FIG. 1), and/or an end effector 300 of the surgical instrument 10. In some examples, the magnetic angle rotary position encoders 2040a-2040b may be coupled to the safety processor 2004 and/or the primary processor 2006.

[0063] In some aspects, the segmented circuit 2000 comprises a motor circuit segment 2002g (Segment 7). The motor circuit segment 2002g comprises a motor 2048 configured to control one or more movements of the powered surgical instrument 10. The motor 2048 is coupled to the primary processor 2006 by an H-Bridge driver 2042 and one or more H-bridge field-effect transistors 2044 (FETs). The H-bridge FETs 2044 are coupled to the safety processor 2004. A motor current sensor 2046 is coupled in series with the motor 2048 to measure the current draw of the motor 2048. The motor current sensor 2046 is in signal communication with the primary processor 2006 and/or the safety processor 2004. In some examples, the motor 2048 is coupled to a motor electromagnetic interference (EMI) filter 2050.

[0064] In some aspects, the segmented circuit 2000 comprises a power segment 2002h (Segment 8). A battery 2008 is coupled to the safety processor 2004, the primary processor 2006, and one or more of the additional circuit segments 2002c-2002g. The battery 2008 is coupled to the segmented circuit 2000 by a battery connector 2010 and a current sensor 2012. The current sensor 2012 is configured to measure the total current draw of the segmented circuit 2000. In some examples, one or more voltage converters 2014a, 2014b, 2016 are configured to provide predetermined voltage values to one or more circuit segments 2002a-2002g. For example, in some examples, the segmented circuit 2000 may comprise 3.3V voltage converters 2014a-2014b and/or 5V voltage converters 2016. A boost converter 2018 is configured to provide a boost voltage up to a predetermined amount, such as, for example, up to 13V. The boost converter 2018 is configured to provide additional voltage and/or current during power intensive operations and prevent brownout or low-power conditions.

[0065] In some aspects, the safety processor segment 2002a comprises a motor power switch 2020. The motor power switch 2020 is coupled between the power segment 2002h and the motor circuit segment 2002g. The safety processor segment 2002a is configured to inter-

rupt power to the motor circuit segment 2002g when an error or fault condition is detected by the safety processor 2004 and/or the primary processor 2006 as discussed in more detail herein. Although the circuit segments 2002a-2002g are illustrated with all components of the circuit segments 2002a-2002h located in physical proximity, one skilled in the art will recognize that a circuit segment 2002a-2002h may comprise components physically and/or electrically separate from other components of

¹⁰ the same circuit segment 2002a-2002g. In some examples, one or more components may be shared between two or more circuit segments 2002a-2002g.

[0066] In some aspects, a plurality of switches 2056-2070 are coupled to the safety processor 2004
 ¹⁵ and/or the primary processor 2006. The plurality of switches 2056-2070 may be configured to control one or more operations of the surgical instrument 10, control one or more operations of the segmented circuit 2000, and/or indicate a status of the surgical instrument 10. For

20 example, a bail-out door switch 2056 is configured to indicate the status of a bail-out door. A plurality of articulation switches, such as, for example, a left side articulation left switch 2058a, a left side articulation right switch 2060a, a left side articulation center switch 2062a,

²⁵ a right side articulation left switch 2058b, a right side articulation right switch 2060b, and a right side articulation center switch 2062b are configured to control articulation of a shaft assembly 200 and/or an end effector 300. A left side reverse switch 2064a and a right side reverse

³⁰ switch 2064b are coupled to the primary processor 2006. In some examples, the left side switches comprising the left side articulation left switch 2058a, the left side articulation right switch 2060a, the left side articulation center switch 2062a, and the left side reverse switch 2064a are

³⁵ coupled to the primary processor 2006 by a left flex connector 2072a. The right side switches comprising the right side articulation left switch 2058b, the right side articulation right switch 2060b, the right side articulation center switch 2062b, and the right side reverse switch

40 2064b are coupled to the primary processor 2006 by a right flex connector 2072b. In some examples, a firing switch 2066, a clamp release switch 2068, and a shaft engaged switch 2070 are coupled to the primary processor 2006.

45 [0067] In some aspects, the plurality of switches 2056-2070 may comprise, for example, a plurality of handle controls mounted to a handle of the surgical instrument 10, a plurality of indicator switches, and/or any combination thereof. In various examples, the plurality of 50 switches 2056-2070 allow a surgeon to manipulate the surgical instrument, provide feedback to the segmented circuit 2000 regarding the position and/or operation of the surgical instrument, and/or indicate unsafe operation of the surgical instrument 10. In some examples, addi-55 tional or fewer switches may be coupled to the segmented circuit 2000, one or more of the switches 2056-2070 may be combined into a single switch, and/or expanded to multiple switches. For example, in one example, one

or more of the left side and/or right side articulation switches 2058a-2064b may be combined into a single multi-position switch.

[0068] In one aspect, the safety processor 2004 is configured to implement a watchdog function, among other safety operations. The safety processor 2004 and the primary processor 2006 of the segmented circuit 2000 are in signal communication. A processor alive heartbeat signal is provided at output 2097. The acceleration segment 2002c comprises an accelerometer 2022 configured to monitor movement of the surgical instrument 10. In various examples, the accelerometer 2022 may be a single, double, or triple axis accelerometer. The accelerometer 2022 may be employed to measures proper acceleration that is not necessarily the coordinate acceleration (rate of change of velocity). Instead, the accelerometer sees the acceleration associated with the phenomenon of weight experienced by a test mass at rest in the frame of reference of the accelerometer 2022. For example, the accelerometer 2022 at rest on the surface of the earth will measure an acceleration g=9.8 m/s² (gravity) straight upwards, due to its weight. Another type of acceleration that accelerometer 2022 can measure is g-force acceleration. In various other examples, the accelerometer 2022 may comprise a single, double, or triple axis accelerometer. Further, the acceleration segment 2002c may comprise one or more inertial sensors to detect and measure acceleration, tilt, shock, vibration, rotation, and multiple degrees-of-freedom (DoF). A suitable inertial sensor may comprise an accelerometer (single, double, or triple axis), a magnetometer to measure a magnetic field in space such as the earth's magnetic field, and/or a gyroscope to measure angular velocity.

[0069] In one aspect, the safety processor 2004 is configured to implement a watchdog function with respect to one or more circuit segments 2002c-2002h, such as, for example, the motor circuit segment 2002g. In this regards, the safety processor 2004 employs the watchdog function to detect and recover from malfunctions of the primary processor 2006. During normal operation, the safety processor 2004 monitors for hardware faults or program errors of the primary processor 2006 and to initiate corrective action or actions. The corrective actions may include placing the primary processor 2006 in a safe state and restoring normal system operation. In one example, the safety processor 2004 is coupled to at least a first sensor. The first sensor measures a first property of the surgical instrument 10 (FIGS. 1-4). In some examples, the safety processor 2004 is configured to compare the measured property of the surgical instrument 10 to a predetermined value. For example, in one example, a magnetic angle rotary position encoder 2040a is coupled to the safety processor 2004. The magnetic angle rotary position encoder 2040a provides motor speed and position information to the safety processor 2004. The safety processor 2004 monitors the magnetic angle rotary position encoder 2040a and compares the value to a maximum speed and/or position value and prevents operation of the motor 2048 above the predetermined values. In some examples, the predetermined values are calculated based on real-time speed and/or position of the motor 2048, calculated from values supplied by a second magnetic angle rotary position encoder 2040b in communication with the primary processor 2006, and/or provided to the safety processor 2004 from, for example, a memory module coupled to the safety processor 2004. **[0070]** In some aspects, a second sensor is coupled

¹⁰ to the primary processor 2006. The second sensor is configured to measure the first physical property. The safety processor 2004 and the primary processor 2006 are configured to provide a signal indicative of the value of the first sensor and the second sensor respectively.

¹⁵ When either the safety processor 2004 or the primary processor 2006 indicates a value outside of an acceptable range, the segmented circuit 2000 prevents operation of at least one of the circuit segments 2002c-2002h, such as, for example, the motor circuit segment 2002g.

For example, in the example illustrated in FIGS. 16A and 16B, the safety processor 2004 is coupled to a first magnetic angle rotary position encoder 2040a and the primary processor 2006 is coupled to a second magnetic angle rotary position encoder 2040b. The magnetic angle rotary

position encoders 2040a, 2040b may comprise any suitable motor position sensor, such as, for example, a magnetic angle rotary input comprising a sine and cosine output. The magnetic angle rotary position encoders 2040a, 2040b provide respective signals to the safety processor
 2004 and the primary processor 2006 indicative of the

position of the motor 2048.

[0071] The safety processor 2004 and the primary processor 2006 generate an activation signal when the values of the first magnetic angle rotary position encoder
 2040a and the second magnetic angle rotary position encoder 2040b are within a predetermined range. When either the primary processor 2006 or the safety processor 2004 to detect a value outside of the predetermined range, the activation signal is terminated and operation
 of at least one of the circuit segments 2002c-2002h, such

as, for example, the motor circuit segment 2002g, is interrupted and/or prevented. For example, in some examples, the activation signal from the primary processor 2006 and the activation signal from the safety processor

45 2004 are coupled to an AND gate. The AND gate is coupled to a motor power switch 2020. The AND gate maintains the motor power switch 2020 in a closed, or on, position when the activation signal from both the safety processor 2004 and the primary processor 2006 are high, 50 indicating a value of the magnetic angle rotary position encoders 2040a, 2040b within the predetermined range. When either of the magnetic angle rotary position encoders 2040a, 2040b detect a value outside of the predetermined range, the activation signal from that magnetic an-55 gle rotary position encoder 2040a, 2040b is set low, and the output of the AND gate is set low, opening the motor power switch 2020. In some examples, the value of the first magnetic angle rotary position encoder 2040a and

the second magnetic angle rotary position encoder 2040b is compared, for example, by the safety processor 2004 and/or the primary processor 2006. When the values of the first sensor and the second sensor are different, the safety processor 2004 and/or the primary processor 2006 may prevent operation of the motor circuit segment 2002g.

[0072] In some aspects, the safety processor 2004 receives a signal indicative of the value of the second magnetic angle rotary position encoder 2040b and compares the second sensor value to the first sensor value. For example, in one aspect, the safety processor 2004 is coupled directly to a first magnetic angle rotary position encoder 2040a. A second magnetic angle rotary position encoder 2040b is coupled to a primary processor 2006, which provides the second magnetic angle rotary position encoder 2040b value to the safety processor 2004, and/or coupled directly to the safety processor 2004. The safety processor 2004 compares the value of the first magnetic angle rotary position encoder 2040 to the value of the second magnetic angle rotary position encoder 2040b. When the safety processor 2004 detects a mismatch between the first magnetic angle rotary position encoder 2040a and the second magnetic angle rotary position encoder 2040b, the safety processor 2004 may interrupt operation of the motor circuit segment 2002g, for example, by cutting power to the motor circuit segment 2002g.

[0073] In some aspects, the safety processor 2004 and/or the primary processor 2006 is coupled to a first magnetic angle rotary position encoder 2040a configured to measure a first property of a surgical instrument and a second magnetic angle rotary position encoder 2040b configured to measure a second property of the surgical instrument. The first property and the second property comprise a predetermined relationship when the surgical instrument is operating normally. The safety processor 2004 monitors the first property and the second property. When a value of the first property and/or the second property inconsistent with the predetermined relationship is detected, a fault occurs. When a fault occurs, the safety processor 2004 takes at least one action, such as, for example, preventing operation of at least one of the circuit segments, executing a predetermined operation, and/or resetting the primary processor 2006. For example, the safety processor 2004 may open the motor power switch 2020 to cut power to the motor circuit segment 2002g when a fault is detected.

[0074] In one aspect, the safety processor 2004 is configured to execute an independent control algorithm. In operation, the safety processor 2004 monitors the segmented circuit 2000 and is configured to control and/or override signals from other circuit components, such as, for example, the primary processor 2006, independently. The safety processor 2004 may execute a preprogrammed algorithm and/or may be updated or programmed on the fly during operation based on one or more actions and/or positions of the surgical instrument 10. For example, in one example, the safety processor 2004 is reprogrammed with new parameters and/or safety algorithms each time a new shaft and/or end effector is coupled to the surgical instrument 10. In some examples, one or more safety values stored by the safety processor 2004 are duplicated by the primary processor 2006. Two-way error detection is performed to ensure values and/or parameters stored by either of the safety processor 2006 are correct.

10 [0075] In some aspects, the safety processor 2004 and the primary processor 2006 implement a redundant safety check. The safety processor 2004 and the primary processor 2006 provide periodic signals indicating normal operation. For example, during operation, the safety

¹⁵ processor 2004 may indicate to the primary processor 2006 that the safety processor 2004 is executing code and operating normally. The primary processor 2006 may, likewise, indicate to the safety processor 2004 that the primary processor 2006 is executing code and oper-

20 ating normally. In some examples, communication between the safety processor 2004 and the primary processor 2006 occurs at a predetermined interval. The predetermined interval may be constant or may be variable based on the circuit state and/or operation of the surgical 25 instrument 10.

[0076] FIGS. 17A and 17B illustrate another aspect of a segmented circuit 3000 configured to control the powered surgical instrument 10, illustrated in FIGS. 1-14. As shown in FIGS. 14, 17B, the handle assembly 14 may
³⁰ include an electric motor 3014 which can be controlled by a motor driver 3015 and can be employed by the firing system of the surgical instrument 10. In various forms, the electric motor 3014 may be a DC brushed driving motor having a maximum rotation of, approximately,
³⁵ 25,000 RPM, for example. In other arrangements, the electric motor 3014 may include a brushless motor, a cordless motor, a synchronous motor, a stepper motor, or any other suitable electric motor. In certain circum-

stances, the motor driver 3015 may comprise an HBridge FETs 3019, as illustrated in FIGS. 17A and 17B, for example. The electric motor 3014 can be powered by a power assembly 3006, which can be releasably mounted to the handle assembly 14. The power assembly 3006 is configured to supply control power to the surgical in-

strument 10. The power assembly 3006 may comprise a battery which may include a number of battery cells connected in series that can be used as the power source to power the surgical instrument 10. In such configuration, the power assembly 3006 may be referred to as a
battery pack. In certain circumstances, the battery cells of the power assembly 3006 may be replaceable and/or rechargeable. In at least one example, the battery cells can be Lithium-Ion batteries which can be separably couplable to the power assembly 3006.

⁵⁵ [0077] Examples of drive systems and closure systems that are suitable for use with the surgical instrument
 10 are disclosed in U.S. Patent Application Publication
 No. 2014/0263539, entitled CONTROL SYSTEMS FOR

SURGICAL INSTRUMENTS, which is incorporated herein by reference herein in its entirety. For example, the electric motor 3014 can include a rotatable shaft (not shown) that may operably interface with a gear reducer assembly that can be mounted in meshing engagement with a set, or rack, of drive teeth on a longitudinally-movable drive member. In use, a voltage polarity provided by the battery can operate the electric motor 3014 to drive the longitudinally-movable drive member to effectuate the end effector 300. For example, the electric motor 3014 can be configured to drive the longitudinally-movable drive member to advance a firing mechanism to fire staples into tissue captured by the end effector 300 from a staple cartridge assembled with the end effector 300 and/or advance a cutting member to cut tissue captured by the end effector 300, for example.

[0078] As illustrated in FIGS. 17A and 17B and as described below in greater detail, the power assembly 3006 may include a power management controller which can be configured to modulate the power output of the power assembly 3006 to deliver a first power output to power the electric motor 3014 to advance the cutting member while the interchangeable shaft assembly 200 is coupled to the handle assembly 14 (FIG. 1) and to deliver a second power output to power the electric motor 3014 to advance the cutting member while the interchangeable shaft assembly 200 is coupled to the handle assembly 14, for example. Such modulation can be beneficial in avoiding transmission of excessive power to the electric motor 3014 beyond the requirements of an interchangeable shaft assembly that is coupled to the handle assembly 14.

[0079] In certain circumstances, the interface 3024 can facilitate transmission of the one or more communication signals between the power management controller 3016 and the shaft assembly controller 3022 by routing such communication signals through a main controller 3017 residing in the handle assembly 14 (FIG. 1), for example. In other circumstances, the interface 3024 can facilitate a direct line of communication between the power management controller 3016 and the shaft assembly controller 3022 through the handle assembly 14 while the interchangeable shaft assembly 200 (FIG. 1) and the power assembly 3006 are coupled to the handle assembly 14. [0080] In one instance, the main controller 3017 may be any single core or multicore processor such as those known under the trade name ARM Cortex by Texas Instruments. In one instance, the surgical instrument 10 (FIGS. 1-4) may comprise a power management controller 3016 such as, for example, a safety controller platform comprising two controller-based families such as TMS570 and RM4x known under the trade name Hercules ARM Cortex R4, also by Texas Instruments. Nevertheless, other suitable substitutes for controllers and safety processor may be employed, without limitation. In one instance, the safety processor 2004 (FIG. 16a) may be configured specifically for IEC 61508 and ISO 26262 safety critical applications, among others, to provide advanced integrated safety features while delivering scalable performance, connectivity, and memory options. **[0081]** In certain instances, the main controller 3017 may be a single core or multicore controller LM4F230H5QR as described in connection with FIGS. 15-17B.

[0082] FIG. 18 is a block diagram the surgical instrument of FIG. 1 illustrating interfaces between the handle assembly 14 (FIG. 1) and the power assembly and be-

tween the handle assembly 14 and the interchangeable shaft assembly. As shown in FIG. 18, the power assembly 3006 may include a power management circuit 3034 which may comprise the power management controller 3016, a power modulator 3038, and a current sense cir-

¹⁵ cuit 3036. The power management circuit 3034 can be configured to modulate power output of the battery 3007 based on the power requirements of the interchangeable shaft assembly 200 (FIG. 1) while the interchangeable shaft assembly 200 and the power assembly 3006 are

²⁰ coupled to the handle assembly 14. For example, the power management controller 3016 can be programmed to control the power modulator 3038 of the power output of the power assembly 3006 and the current sense circuit 3036 can be employed to monitor power output of the

²⁵ power assembly 3006 to provide feedback to the power management controller 3016 about the power output of the battery 3007 so that the power management controller 3016 may adjust the power output of the power assembly 3006 to maintain a desired output.

30 [0083] It is noteworthy that the power management controller 3016 and/or the shaft assembly controller 3022 each may comprise one or more processors and/or memory units which may store a number of software modules. Although certain modules and/or blocks of the surgical

instrument 10 (FIG. 1) may be described by way of example, it can be appreciated that a greater or lesser number of modules and/or blocks may be used. Further, although various instances may be described in terms of modules and/or blocks to facilitate description, such mod-

 ⁴⁰ ules and/or blocks may be implemented by one or more hardware components, e.g., processors, Digital Signal Processors (DSPs), Programmable Logic Devices (PLDs), Application Specific Integrated Circuits (ASICs), circuits, registers and/or software components, e.g., pro ⁴⁵ grams, subroutines, logic and/or combinations of hard-

grams, subroutines, logic and/or combinations of hardware and software components.

[0084] In certain instances, the surgical instrument 10 (FIGS. 1-4) may comprise an output device 3042 which may include one or more devices for providing a sensory
feedback to a user. Such devices may comprise, for example, visual feedback devices (e.g., an LCD display screen, LED indicators), audio feedback devices (e.g., a speaker, a buzzer) or tactile feedback devices (e.g., haptic actuators). In certain circumstances, the output device
3042 may comprise a display 3043 which may be included in the handle assembly 14 (FIG. 1). The shaft assembly controller 3022 and/or the power management controller 3016 can provide feedback to a user of the surgical

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instrument 10 through the output device 3042. The interface 3024 can be configured to connect the shaft assembly controller 3022 and/or the power management controller 3016 to the output device 3042. The reader will appreciate that the output device 3042 can instead be integrated with the power assembly 3006. In such circumstances, communication between the output device 3042 and the shaft assembly controller 3022 may be accomplished through the interface 3024 while the interchangeable shaft assembly 200 is coupled to the handle assembly 14.

[0085] Having described a surgical instrument 10 (FIGS. 1-4) and one or more segmented circuit 2000, 3000 for controlling the operation thereof, the disclosure now turns to various specific configurations of the surgical instrument 10 and a segmented circuit 2000 (or 3000). [0086] In various aspects the present disclosure provides techniques for data storage and usage. In one aspect, data storage and usage is based on multiple levels of action thresholds. Such thresholds include upper and lower ultimate threshold limits, ultimate threshold that shuts down motor or activates return is current, pressure, firing load, torque is exceeded, and alternatively, while running within the limits the device automatically compensates for loading of the motor.

[0087] In one aspect, the surgical instrument 10 (described in connection with FIGS. 1-18) can be configured to monitor upper and lower ultimate threshold limits to maintain minimum and maximum closure clamp loads within acceptable limits. If a minimum is not achieved the surgical instrument 10 cannot start or if it drops below minimum a user action is required. If the clamp load is at a suitable level but drops under minimum during firing, the surgical instrument 10 can adjust the speed of the motor or warn the user. If the minimum limit is breached during operation the unit could give a warning that the firing may not be completely as anticipated. The surgical instrument 10 also can be configured to monitor when the battery voltage drops below the lower ultimate limit the remaining battery power is only direct able towards returning the device to the I-beam parked state. The opening force on the anvil can be employed to sense jams in the end effector. Alternatively, the surgical instrument 10 can be configured to monitor when the motor current goes up or the related speed goes down, then the motor control increases pulse width or frequency modulation to keep speed constant.

[0088] In another aspect, the surgical instrument 10 can (FIG. 1) be configured to detect an ultimate threshold of current draw, pressure, firing load, torque such that when any of these thresholds are exceeded, the surgical instrument 10 shuts down the motor or causes the motor to return the knife to a pre-fired position. A secondary threshold, which is less than the ultimate threshold, may be employed to alter the motor control program to accommodate changes in conditions by changing the motor control parameters. A marginal threshold can be configured as a step function or a ramp function based on a

proportionate response to another counter or input. For example, in the case of sterilization, no changes between 0-200 sterilization cycles, slow motor 1% per use from 201-400 sterilization cycles, and prevent use over 400 sterilization cycles. The speed of the motor also can be varied based on tissue gap and current draw.

[0089] There are many parameters that could influence the ideal function of a powered reusable stapler device. Most of these parameters have an ultimate max-

¹⁰ imum and/or minimum threshold beyond which the device should not be operated. Nevertheless, there are also marginal limits that may influence the functional operation of the device. These multiple limits, from multiple parameters may provide an overlying and cumulative ef-¹⁵ fect on the operations program of the device.

[0090] Accordingly, the present disclosure relates to surgical instruments and, in various circumstances, to surgical stapling and cutting instruments and staple cartridges therefor that are designed to staple and cut tissue.

20 [0091] Efficient performance of an electromechanical device depends on various factors. One is the operational envelope, i.e., range of parameters, conditions and events in which the device carries out its intended functions. For example, for a device powered by a motor driv-

en by electrical current, there may be an operational region above a certain electrical current threshold where the device runs more inefficiently than desired. Put another way, there may be an upper "speed limit" above which there is decreasing efficiency. Such an upper
threshold may have value in preventing substantial inefficiencies or even device degradation.

[0092] There may be thresholds within an operational envelope, however, that may form regions exploitable to enhance efficiency within operational states. In other ³⁵ words, there may be regions where the device can adjust and perform better within a defined operational envelope (or sub-envelope). Such a region can be one between a marginal threshold and an ultimate threshold. In addition, these regions may comprise "sweet spots" or a prede-

40 termined optional range or point. These regions also may comprise a large range within which performance is judged to be adequate.

[0093] An ultimate threshold can be defined, above which or below which an action or actions could be taken

- ⁴⁵ (or refrained from being taken) such as stopping the device. In addition, a marginal threshold or thresholds can be defined, above which or below which an action or actions could be taken (or refrained from being taken). By way of non-limiting example, a marginal threshold can
- ⁵⁰ be set to define where the current draw of the motor exceeds 75% of an ultimate threshold. Exceeding the marginal threshold can result, for example, in the device's beginning to slow motor speed at an increasing rate as it continues to climb toward the ultimate threshold.

⁵⁵ **[0094]** Various mechanisms can be employed to carry out the adjustment(s) taken as a result of exceeding a threshold. For example, the adjustment can reflect a step function. It can also reflect a ramped function. Other func-

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tions can be utilized.

[0095] In various aspects, to enhance performance by additional mechanisms, an overlaying threshold can be defined. An overlaying threshold can comprise one or more thresholds defined by multiple parameters. An overlaying threshold can result in one or more thresholds being an input into the generation of another threshold or thresholds. An overlaying threshold can be predetermined or dynamically generated such as at runtime. The overlaying threshold may come into effect when you the threshold is defined by multiple inputs. For example, as the number of sterilization cycles exceeds 300 (the marginal threshold) but not 500 (the ultimate threshold) the device runs the motor slower. Then as the current draw exceeds its 75% marginal threshold it multiples the slow down going even slower.

[0096] FIG. 19 illustrates a logic diagram of a system 4311 for evaluating sharpness of a cutting edge 182 (FIG. 14) of a surgical instrument 10 (FIGS. 1-4) according to various examples. In certain instances, the system 4311 can evaluate the sharpness of the cutting edge 182 by testing the ability of the cutting edge 182 to be advanced through a sharpness testing member 4302. For example, the system 4311 can be configured to observe the time period the cutting edge 182 takes to fully transect and/or completely pass through at least a predetermined portion of a sharpness testing member 4302. If the observed time period exceeds a predetermined threshold, the circuit 4310 may conclude that the sharpness of the cutting edge 182 has dropped below an acceptable level, for example.

[0097] In one aspect, the sharpness testing member 4302 can be employed to test the sharpness of the cutting edge 182 (FIG. 14). In certain instances, the sharpness testing member 4302 can be attached to and/or integrated with the cartridge body 194 (FIG. 14) of the surgical staple cartridge 304 (FIGS. 1, 2, and 15), for example. In certain instances, the sharpness testing member 4302 can be disposed in the proximal portion of the surgical staple cartridge 304, for example. In certain instances, the sharpness testing member 4302 can be disposed onto a cartridge deck or cartridge body 194 of the surgical staple cartridge 304, for example.

[0098] In certain instances, a load cell 4335 can be configured to monitor the force (Fx) applied to the cutting edge 182 (FIG. 14) while the cutting edge 182 is engaged and/or in contact with the sharpness testing member 4302, for example. The reader will appreciate that the force (Fx) applied by the sharpness testing member 4302 to the cutting edge 182 while the cutting edge 182 is engaged and/or in contact with the sharpness testing member 4302 to the cutting edge 182 while the cutting edge 182 is engaged and/or in contact with the sharpness testing member 4302 may depend, at least in part, on the sharpness of the cutting edge 182. In certain instances, a decrease in the sharpness of the cutting edge 182 can result in an increase in the force (Fx) required for the cutting edge 182 to cut or pass through the sharpness testing member 4302. The load cell 4335 of the sharpness testing member 4302 may be employed to measure the force

(Fx) applied to the cutting edge 182 while the cutting edge 182 travels a predefined distance (D) through the sharpness testing member 4302 may be employed to determine the sharpness of the cutting edge 182.

⁵ [0099] In certain instances, the system 4311 may include a controller 4313 ("microcontroller") which may include a processor 4315 ("microprocessor") and one or more computer readable mediums or memory 4317 units ("memory"). In certain instances, the memory 4317 may

¹⁰ store various program instructions, which when executed may cause the processor 4315 to perform a plurality of functions and/or calculations described herein. In certain instances, the memory 4317 may be coupled to the processor 4315, for example. A power source 4319 can be

¹⁵ configured to supply power to the controller 4313, for example. In certain instances, the power source 4319 may comprise a battery (or "battery pack" or "power pack"), such as a Li ion battery, for example. In certain instances, the battery pack may be configured to be re-

²⁰ leasably mounted to the handle assembly 14. A number of battery cells connected in series may be used as the power source 4319. In certain instances, the power source 4319 may be replaceable and/or rechargeable, for example.

²⁵ **[0100]** In certain instances, the controller 4313 can be operably coupled to the feedback system and/or the lock-out mechanism 4123, for example.

[0101] The system 4311 may comprise one or more position sensors. Example position sensors and positioning systems suitable for use with the present disclosure are described in U.S. Patent Application Publication No. 2014/0263538, entitled SENSOR ARRANGEMENTS FOR ABSOLUTE POSITIONING SYSTEM FOR SUR-GICAL INSTRUMENTS, which is herein incorporated by reference in its entirety. In certain instances, the system

4311 may include a first position sensor 4321 and a second position sensor 4323. In certain instances, the first position sensor 4321 can be employed to detect a first position of the cutting edge 182 (FIG. 14) at a proximal
end of a sharpness testing member 4302, for example;

and the second position sensor 4323 can be employed to detect a second position of the cutting edge 182 at a distal end of a sharpness testing member 4302, for example.

45 In certain instances, the first and second posi-[0102] tion sensors 4321, 4323 can be employed to provide first and second position signals, respectively, to the controller 4313. It will be appreciated that the position signals may be analog signals or digital values based on the 50 interface between the controller 4313 and the first and second position sensors 4321, 4323. In one example, the interface between the controller 4313 and the first and second position sensors 4321, 4323 can be a standard serial peripheral interface (SPI), and the position sig-55 nals can be digital values representing the first and second positions of the cutting edge 182, as described above.

[0103] Further to the above, the processor 4315 may

determine the time period between receiving the first position signal and receiving the second position signal. The determined time period may correspond to the time it takes the cutting edge 182 (FIG. 14) to advance through a sharpness testing member 4302 from the first position at a proximal end of the sharpness testing member 4302, for example, to a second position at a distal end of the sharpness testing member 4302, for example. In at least one example, the controller 4313 may include a time element which can be activated by the processor 4315 upon receipt of the first position signal, and deactivated upon receipt of the second position signal. The time period between the activation and deactivation of the time element may correspond to the time it takes the cutting edge 182 to advance from the first position to the second position, for example. The time element may comprise a real time clock, a processor configured to implement a time function, or any other suitable timing circuit.

[0104] In various instances, the controller 4313 can compare the time period it takes the cutting edge 182 (FIG. 14) to advance from the first position to the second position to a predefined threshold value to assess whether the sharpness of the cutting edge 182 has dropped below an acceptable level, for example. In certain instances, the controller 4313 may conclude that the sharpness of the cutting edge 182 has dropped below an acceptable level if the measured time period exceeds the predefined threshold value by 1%, 5%, 10%, 25%, 50%, 100% and/or more than 100%, for example.

[0105] FIG. 20 illustrates a logic diagram of a system 4340 for determining the forces applied against a cutting edge of a surgical instrument 10 (FIGS. 1-4) by a sharpness testing member 4302 at various sharpness levels according to various aspects. Referring to FIG. 20, in various instances, an electric motor 4331 can drive the firing bar 172 (FIG. 20) to advance the cutting edge 182 (FIG. 14) during a firing stroke and/or to retract the cutting edge 182 during a return stroke, for example. A motor driver 4333 can control the electric motor 4331; and a controller such as, for example, the controller 4313 can be in signal communication with the motor driver 4333. As the electric motor 4331 advances the cutting edge 182, the controller 4313 can determine the current drawn by the electric motor 4331, for example. In such instances, the force required to advance the cutting edge 182 can correspond to the current drawn by the electric motor 4331, for example. Referring still to FIG. 20, the controller 4313 of the surgical instrument 10 can determine if the current drawn by the electric motor 4331 increases during advancement of the cutting edge 182 and, if so, can calculate the percentage increase of the current.

[0106] In certain instances, the current drawn by the electric motor 4331 may increase significantly while the cutting edge 182 (FIG. 14) is in contact with the sharpness testing member 4302 due to the resistance of the sharpness testing member 4302 to the cutting edge 182. For example, the current drawn by the electric motor 4331 may increase significantly as the cutting edge 182 en-

gages, passes and/or cuts through the sharpness testing member 4302. The reader will appreciate that the resistance of the sharpness testing member 4302 to the cutting edge 182 depends, in part, on the sharpness of the cutting edge 182; and as the sharpness of the cutting edge

182 decreases from repetitive use, the resistance of the sharpness testing member 4302 to the cutting edge 182 will increase. Accordingly, the value of the percentage increase of the current drawn by the electric motor 4331

¹⁰ while the cutting edge is in contact with the sharpness testing member 4302 can increase as the sharpness of the cutting edge 182 decreases from repetitive use, for example.

[0107] In certain instances, the determined value of
the percentage increase of the current drawn by the electric motor 4331 can be the maximum detected percentage increase of the current drawn by the electric motor
4331. In various instances, the controller 4313 can compare the determined value of the percentage increase of
the current drawn by the electric motor 4331 to a predefined threshold value of the percentage increase of the current drawn by the electric motor 4331. If the determined value exceeds the predefined threshold value, the controller 4313 may conclude that the sharpness of the

⁵ cutting edge 182 has dropped below an acceptable level, for example.

[0108] In certain instances, as illustrated in FIG. 20, the processor 4315 can be in communication with the feedback system and/or the lockout mechanism for ex-30 ample. In certain instances, the processor 4315 can employ the feedback system to alert a user if the determined value of the percentage increase of the current drawn by the electric motor 4331 exceeds the predefined threshold value, for example. In certain instances, the processor 35 4315 may employ the lockout mechanism to prevent advancement of the cutting edge 182 (FIG. 14) if the determined value of the percentage increase of the current drawn by the electric motor 4331 exceeds the predefined threshold value, for example. In certain instances, the 40 system 4311 may include first and second position sen-

sors 4321, 4323. The surgical instrument 10 (FIGS. 1-4) may include a load cell 4335.

[0109] In various instances, the controller 4313 can utilize an algorithm to determine the change in current

45 drawn by the electric motor 4331. For example, a current sensor can detect the current drawn by the electric motor 4331 during the firing stroke. The current sensor can continually detect the current drawn by the electric motor and/or can intermittently detect the current draw by the 50 electric motor. In various instances, the algorithm can compare the most recent current reading to the immediately proceeding current reading, for example. Additionally or alternatively, the algorithm can compare a sample reading within a time period X to a previous current read-55 ing. For example, the algorithm can compare the sample reading to a previous sample reading within a previous time period X, such as the immediately proceeding time

period X, for example. In other instances, the algorithm

can calculate the trending average of current drawn by the motor. The algorithm can calculate the average current draw during a time period X that includes the most recent current reading, for example, and can compare that average current draw to the average current draw during an immediately proceeding time period time X, for example.

[0110] In certain instances, the load cell 4335 (FIGS. 19,20) can be configured to monitor the force (Fx) applied to the cutting edge 182 (FIG. 14) while the cutting edge 182 is engaged and/or in contact with the sharpness testing member 4302 (FIGS. 19, 20), for example. The reader will appreciate that the force (Fx) applied by the sharpness testing member 4302 to the cutting edge 182 while the cutting edge 182 is engaged and/or in contact with the sharpness testing member 4302 may depend, at least in part, on the sharpness of the cutting edge 182. In certain instances, a decrease in the sharpness of the cutting edge 182 can result in an increase in the force (Fx) required for the cutting edge 182 to cut or pass through the sharpness testing member 4302. In certain instances, the controller 4313 (FIGS. 19, 20) may compare a maximum value of the monitored force (Fx) applied to the cutting edge 182 (FIG. 14) to one or more predefined threshold values.

[0111] In certain instances, the cutting edge 182 (FIG. 14) may be sufficiently sharp for transecting a captured tissue comprising a first thickness but may not be sufficiently sharp for transecting a captured tissue comprising a second thickness greater than the first thickness, for example. In certain instances, a sharpness level of the cutting edge 182, as defined by the force required for the cutting edge 182 to transect a captured tissue, may be adequate for transecting the captured tissue if the captured tissue comprises a tissue thickness that is in a particular range of tissue thicknesses, for example. In certain instances, the memory 4317 (FIGS. 19, 20) can store one or more predefined ranges of tissue thicknesses of tissue captured by the end effector 300; and predefined threshold forces associated with the predefined ranges of tissue thicknesses. In certain instances, each predefined threshold force may represent a minimum sharpness level of the cutting edge 182 that is suitable for transecting a captured tissue comprising a tissue thickness (Tx) encompassed by the range of tissue thicknesses that is associated with the predefined threshold force. In certain instances, when the force (Fx) required for the cutting edge 182 to transect the captured tissue, comprising the tissue thickness (Tx), exceeds the predefined threshold force associated with the predefined range of tissue thicknesses that encompasses the tissue thickness (Tx), the cutting edge 182 may not be sufficiently sharp to transect the captured tissue, for example.

[0112] In various aspects, the present disclosure provides techniques for determining tissue compression and additional techniques to control the operation of the surgical instrument 10 (described in connection with FIGS. 1-18) in response to the tissue compression. In one ex-

ample, the cartridges may be configured to define variable compression algorithm which drives the surgical instrument 10 to close differently based on intended tissue type and thickness. In another example, the surgical instrument 10 learns from surgeon use and original tissue compression profile to adapt closure based on load experienced during firing. When the surgical instrument 10 experiences tissue compression loads that are dramatically different that those experienced for this cartridge type the instrument highlights this to the user.

¹⁰ type the instrument highlights this to the user. [0113] Active adjustment of a motor control algorithm over time as the instrument become acclimated to the hospital's usage can improve the life expectancy of a rechargeable battery as well as adjust to tissue/proce-¹⁵ dure requirements of minimizing tissue flow, thus improv-

ing staple formation in the tissue seal.

[0114] Accordingly, the present disclosure relates to surgical instruments and, in various circumstances, to surgical stapling and cutting instruments and staple car tridges therefor that are designed to staple and cut tissue. For example, in various aspects the present disclosure provides an endosurgical instrument configured to sense the cartridge type or tissue gap to enable the handle to adjust the closure and firing algorithms to adjust for in-

tended tissue properties. This adaptive algorithm adjustment can "learn" from the user's operations allowing the device to react and benefit two different systems. The first benefit provided by the disclosed adaptive algorithm includes tissue flow and staple formation. As the device
learns the users' basic habits and step timings, the device can adjust the closure speed and firing speed to provide

a more consistent and reliable output. The second benefit provided by the disclosed adaptive algorithm is related to the battery pack. As the device learns how many firings ³⁵ and what conditions the instrument was used, the device

can adjust motor current needs/speed in a predefined manner to prolong battery life. There is a substantially small likelihood that a device used in a hospital that performs predominantly bariatric procedures would be operated in a manner similar to a device used in a hospital

40 erated in a manner similar to a device used in a hospital that performs mostly colorectal or thoracic procedures. Thus, when the device is used to perform substantially similar procedure, over time, the device is configured to learn and adjust its operational algorithm to maintain

⁴⁵ within the "ideal" discharge and tissue flow envelopes.
[0115] Safe and effective surgery requires due knowledge of, and respect for, the tissue involved. Clinicians are mindful that adjustments made during surgery may be beneficial. These adjustments include mechanisms
⁵⁰ to detect and promote desirable staple formation.

[0116] Endosurgical instruments can generate, monitor and process a substantial amount of data during their use in connection with a surgical procedure. Such data can be obtained from the surgical instrument itself, including battery usage. Additionally, data can be obtained from the properties of the tissue with which the surgical instrument interacts, including properties such as tissue compression. Further, data can be obtained from the cli-

nician's interaction with the surgical instrument itself. The repository of data so obtained can be processed and, where desired, the surgical instrument can be designed to adapt to circumstances so as to promote a safe and effective outcome to the current surgical procedure, as well as lay the foundation for more generalized productive use by multiple clinicians. Such adaptive adjustments -- both during a surgical procedure, and wherein the instrument "learns" based on usage patterns drawn from multiple surgical procedures-can provide numerous mechanisms to enhance the overall patient-care environment.

[0117] FIG. 21 illustrates one aspect of a process for adapting operations of a surgical instrument. As depicted in FIG. 21, a module can be attached 5160 or otherwise loaded to the surgical instrument 10 (FIGS. 1-4). The module can contain a program that is selected or uploaded 5162. Controls can be activated 5164 such that they can be ready to operate the surgical instrument 10. During or after usage of the surgical instrument 10, control measures can be included to adapt 5166 a program. For example, this can include adjusting the data rate within the surgical instrument 10 or with respect to remote operation of the surgical instrument 10. This can include adjusting speed, such as speed by which anvil 306 (FIG. 1) and surgical staple cartridge 304 (FIG. 1) engage in a closure motion. This can also include a pulse from an emitter and sensor or to apply a pulse of electrical current to tissue, and the timing of such pulse. This can include adjusting a program to adapt to acceleration, such as acceleration of the surgical instrument 10 if dropped, or transition from a sleep mode. A program can be adapted to handle an actual and/or expected load based on clamping force.

[0118] The surgical instrument 10 (FIGS. 1-4) can be employed to complete an action 5168, for example to carry out a stapling procedure. Data can be recorded 5170 in appropriate memory locations of the surgical instrument 10. Sensor behavior 5172 can be assessed, such as to what extent a sensor accurately measured and/or measures a parameter. Anticipated data can be assessed 5174, including but not limited to tissue properties, wait period and firing speed. Foregoing mechanisms disclosed herein can provide an input to adapt 5166 a program further. In addition, a tissue identification 5178 can be performed, based on historical, actual or expected tissue properties, and this can provide an input to further adapt 5166 a program. In addition, tissue identification 5178 properties can be updated. Moreover, measured sensor input 5176 during a procedure can be used as an additional input to further adapt 5166 a program; such sensor measurements can include those of the gap between anvil 306 and surgical staple cartridge 304, obtaining a derivative measurement including a derivative of a function, current, or torque.

[0119] The end-effector 6006 may be used to compress, cut, or staple tissue. Referring now to FIG. 23A, an end-effector 6030 may be positioned by a physician

to surround tissue 6032 prior to compression, cutting, or stapling. As shown in FIG. 23A, no compression may be applied to the tissue while preparing to use the end-effector. Referring now to FIG. 23B, by engaging the handle

(e.g., handle 6002) of the endocutter, the physician may use the end-effector 6030 to compress the tissue 6032. In one aspect, the tissue 6032 may be compressed to its maximum threshold, as shown in FIG. 23B.

[0120] Referring to FIG. 23A, various forces may be applied to the tissue 6032 by the end-effector 6030. For example, vertical forces F1 and F2 may be applied by the anvil 6034 and the channel frame 6036 of the endeffector 6030 as tissue 6032 is compressed between the two. Referring now to FIG. 23B, various diagonal and/or

¹⁵ lateral forces also may be applied to the tissue 6032 when compressed by the end-effector 6030. For example, force F3 may be applied. For the purposes of operating a medical device such as endocutter 6000, it may be desirable to sense or calculate the various forms of com-

²⁰ pression being applied to the tissue by the end-effector. For example, knowledge of vertical or lateral compression may allow the end-effector to more precisely or accurately apply a staple operation or may inform the operator of the endocutter such that the endocutter can be used more properly or safely.

[0121] The compression through tissue 6032 may be determined from an impedance of tissue 6032. At various levels of compression, the impedance Z of tissue 6032 may increase or decrease. By applying a voltage V and

³⁰ a current I to the tissue 6032, the impedance Z of the tissue 6032 may be determined at various levels of compression. For example, impedance Z may be calculated by dividing the applied voltage V by the current I.

[0122] Referring now to FIG. 24, in one aspect, an RF
 ³⁵ electrode 6038 may be positioned on the end-effector
 6030 (e.g., on a staple cartridge, knife, or channel frame of the end-effector 6030). Further, an electrical contact
 6040 may be positioned on the anvil 6034 of the end-effector 6030. In one aspect, the electrical contact may

⁴⁰ be positioned on the channel frame of the end-effector. As the tissue 6032 is compressed between the anvil 6034 and, for example, the channel frame 6036 of the endeffector 6030, an impedance Z of the tissue 6032 changes. The vertical tissue compression 6042 caused by the

⁴⁵ end-effector 6030 may be measured as a function of the impedance Z of the tissue 6032.

[0123] Referring now to FIG. 25, in one aspect, an electrical contact 6044 may be positioned on an opposite end of the anvil 6034 of the end-effector 6030 as the RF electrode 6038 is positioned. As the tissue 6032 is compressed between the anvil 6034 and, for example, the channel frame 6036 of the end-effector 6030, an impedance Z of the tissue 6032 changes. The lateral tissue compression 6046 caused by the end-effector 6030 may be measured as a function of the impedance Z of the tissue 6032.

[0124] Referring now to FIG. 26, in one aspect, electrical contact 6050 may be positioned on the anvil 6034

and electrical contact 6052 may be positioned on an opposite end of the end-effector 6030 at channel frame 6036. RF electrode 6048 may be positioned laterally to the central to the end-effector 6030. As the tissue 6032 is compressed between the anvil 6034 and, for example, the channel frame 6036 of the end-effector 6030, an impedance Z of the tissue 6032 changes. The lateral compression or angular compressions 6054 and 6056 on either side of the RF electrode 6048 may be caused by the end-effector 6030 and may be measured as a function of different impedances Z of the tissue 6032, based on the relative positioning of the RF electrode 6048 and electrical contacts 6050 and 6052.

[0125] In accordance with one or more of the techniques and features described in the present disclosure, and as discussed above, an RF electrode may be used as an RF sensor. Referring now to FIG. 27, in one aspect, an RF sensor 6062 may be positioned on a staple cartridge 6060 inserted into a channel frame 6066 an endeffector. The RF electrode may run from a power line 6064 which may be powered by a power source in a handle (e.g., handle 6002) of an endocutter.

[0126] Referring now to FIG. 28, in one aspect, RF electrodes 6074 and 6076 may be positioned on a staple cartridge 6072 inserted into a channel frame 6078 of end-effector 6070. As shown, RF electrode 6074 may be placed in a proximal position of the end-effector relative to an endocutter handle. Further, RF electrode 6076 may be placed in a distal position of the end-effector relative to the endocutter handle. RF electrodes 6074 and 6076 may be utilized to measure vertical, lateral, proximal, or distal compression at different points in a tissue based on the position of one or more electrical contacts on the end-effector.

[0127] Referring now to FIG. 29, in one aspect, RF electrodes 6084-6116 may be positioned on staple cartridge 6082 inserted into the channel frame 6080 (or other component of an end-effector) based on various points for which compression information is desired. Referring now to FIG. 30, in one aspect, RF electrodes 6122-6140 may be positioned on staple cartridge 6120 at discrete points for which compression information is desired. Referring now to FIG. 31, RF electrodes 6152-6172 may be positioned at different points in multiple zones of a staple cartridge based on how accurate or precise the compression measurements should be. For example, RF electrodes 6152-6156 may be positioned in zone 6158 of staple cartridge 6150 depending on how accurate or precise the compression measurements in zone 6158 should be. Further, RF electrodes 6160-6164 may be positioned in zone 6166 of staple cartridge 6150 depending on how accurate or precise the compression measurements in zone 6166 should be. Additionally, RF electrodes 6168-6172 may be positioned in zone 6174 of staple cartridge 6150 depending on how accurate or precise the compression measurements in zone 6174 should be.

[0128] The RF electrodes discussed herein may be

wired through a staple cartridge inserted in the channel frame. Referring now to FIG. 32, in one aspect, an RF electrode may have a stamped "mushroom head" 6180 of about 1.0 mm in diameter. While the RF electrode may have the stamped "mushroom head" of about 1.0 mm in

diameter, this is intended to be a non-limiting example and the RF electrode may be differently shaped and sized depending on each particular application or design. The RF electrode may be connected to, fastened to, or may

form, a conductive wire 6182. The conductive wire 6182 may be about 0.5 mm in diameter, or may have a larger or smaller diameter based on a particular application or design. Further, the conductive wire may have an insulative coating 6184. In one example, the RF electrode may protrude through a staple cartridge, channel frame,

knife, or other component of an end-effector.
[0129] Referring now to FIG. 33, the RF electrodes may be wired through a single wall or through multiple walls of a staple cartridge or channel frame of an end-effector. For example, RF electrodes 6190-6194 may be wired through wall 6196 of the staple cartridge or channel frame of an end-effector. One or more of wires 6198 may be connected to, fastened to, or be part of, RF electrodes 6190-6194 and may run through wall 6196 from a power source in, e.g., a handle of an endocutter.

[0130] Referring now to FIG. 34, the power source may be in communication with the RF electrodes or may provide power to the RF electrodes through a wire or cable. The wire or cable may join each individual wire and lead
 30 to the power source. For example, RF electrodes

6204-6212 may receive power from a power source through wire or cable 6202, which may run through staple cartridge 6200 or a channel frame of an end-effector. In one example, each of RF electrodes 6204-6212 may have its own wire that runs to or through wire or cable

³⁵ have its own wire that runs to or through wire or cable 6202. The staple cartridge 6200 or channel frame also may include a controller 6214, such as the primary processor 2006 shown in connection with FIGS. 16A and 16B, or the main controller 3017 shown in connection with

40 FIGS. 17A, 17B, and 18, for example. It will be appreciated that the controller 6214 should be suitably sized to fit in the staple cartridge 6200 or channel frame form factor. Also, the controller

[0131] In various aspects, the tissue compression sen-45 sor system described herein for use with medical devices may include a frequency generator. The frequency generator may be located on a circuit board of the medical device, such as an endocutter. For example the frequency generator may be located on a circuit board in a shaft 50 or handle of the endocutter. Referring now to FIG. 35, an example circuit diagram 6220 in accordance with one example of the present disclosure is shown. As shown, frequency generator 6222 may receive power or current from a power source 6221 and may supply one or more 55 RF signals to one or more RF electrodes 6224. As discussed above, the one or more RF electrodes may be positioned at various locations or components on an endeffector or endocutter, such as a staple cartridge or chan-

nel frame. One or more electrical contacts, such as electrical contacts 6226 or 6228 may be positioned on a channel frame or an anvil of an end-effector. Further, one or more filters, such as filters 6230 or 6232 may be communicatively coupled to the electrical contacts 6226 or 6228 as shown in FIG. 35. The filters 6230 and 6232 may filter one or more RF signals supplied by the frequency generator 6222 before joining a single return path 6234. A voltage V and a current I associated with the one or more RF signals may be used to calculate an impedance Z associated with a tissue that may be compressed and/or communicatively coupled between the one or more RF electrodes 6224 and the electrical contacts 6226 or 6228.

[0132] Referring now to FIG. 36, various components of the tissue compression sensor system described herein may be located in a handle 6236 of an endocutter. For example, as shown in circuit diagram 6220a, frequency generator 6222 may be located in the handle 6236 and receives power from power source 6221. Also, current I1 and current I2 may be measured on a return path corresponding to electrical contacts 6228 and 6226. Using a voltage V applied between the supply and return paths, impedances Z1 and Z2 may be calculated. Z1 may correspond to an impedance of a tissue compressed and/or communicatively coupled between one or more of RF electrodes 6224 and electrical contact 6228. Further, Z2 may correspond to an impedance of a tissue compressed and/or communicatively coupled between one or more of RF electrodes 6224 and electrical contact 6226. Applying the formulas Z1 = V/I1 and Z2 = V/I2, impedances Z1 and Z2 corresponding to different compression levels of a tissue compressed by an end-effector may be calculated.

[0133] Referring now to FIG. 37, one or more aspects of the present disclosure are described in circuit diagram 6250. In an implementation, a power source at a handle 6252 of an endocutter may provide power to a frequency generator 6254. The frequency generator 6254 may generate one or more RF signals. The one or more RF signals may be multiplexed or overlaid at a multiplexer 6256, which may be in a shaft 6258 of the endocutter. In this way, two or more RF signals may be overlaid (or, e.g., nested or modulated together) and transmitted to the end-effector. The one or more RF signals may energize one or more RF electrodes 6260 at an end-effector 6262 (e.g., positioned in a staple cartridge) of the endocutter. A tissue (not shown) may be compressed and/or communicatively coupled between the one or more of RF electrodes 6260 and one or more electrical contacts. For example, the tissue may be compressed and/or communicatively coupled between the one or more RF electrodes 6260 and the electrical contact 6264 positioned in a channel frame of the end-effector 6262 or the electrical contact 6266 positioned in an anvil of the end-effector 6262. A filter 6268 may be communicatively coupled to the electrical contact 6264 and a filter 6270 may be communicatively coupled to the electrical contact

6266.

[0134] A voltage V and a current I associated with the one or more RF signals may be used to calculate an impedance Z associated with a tissue that may be compressed between the staple cartridge (and communicatively coupled to one or more RF electrodes 6260) and the channel frame or anvil (and communicatively coupled to one or more of electrical contacts 6264 or 6266).

[0135] In one aspect, various components of the tissue
 compression sensor system described herein may be located in a shaft 6258 of the endocutter. For example, as shown in circuit diagram 6250 (and in addition to the frequency generator 6254), an impedance calculator 6272, a controller 6274, a non-volatile memory 6276, and

¹⁵ a communication channel 6278 may be located in the shaft 6258. In one example, the frequency generator 6254, impedance calculator 6272, controller 6274, nonvolatile memory 6276, and communication channel 6278 may be positioned on a circuit board in the shaft 6258.

20 [0136] The two or more RF signals may be returned on a common path via the electrical contacts. Further, the two or more RF signals may be filtered prior to the joining of the RF signals on the common path to differentiate separate tissue impedances represented by the

²⁵ two or more RF signals. Current I1 and current I2 may be measured on a return path corresponding to electrical contacts 6264 and 6266. Using a voltage V applied between the supply and return paths, impedances Z1 and Z2 may be calculated. Z1 may correspond to an imped-

ance of a tissue compressed and/or communicatively coupled between one or more of RF electrodes 6260 and electrical contact 6264. Further, Z2 may correspond to an impedance of the tissue compressed and/or communicatively coupled between one or more of RF electrodes
 6260 and electrical contact 6266. Applying the formulas

6260 and electrical contact 6266. Applying the formulas Z1 = V/I1 and Z2 = V/I2, impedances Z1 and Z2 corresponding to different compressions of a tissue compressed by an end-effector 6262 may be calculated. In example, the impedances Z1 and Z2 may be calculated

40 by the impedance calculator 6272. The impedances Z1 and Z2 may be used to calculate various compression levels of the tissue.

[0137] In one aspect, filters 6268 and 6270 may be High Q filters such that the filter range may be narrow

⁴⁵ (e.g., Q=10). Q may be defined by the Center frequency (Wo)/Bandwidth (BW) where Q= Wo/BW. In one example, Frequency 1 may be 150kHz and Frequency 2 may be 300kHz. A viable impedance measurement range may be 100kHz - 20 MHz. In various examples, other sophisticated techniques, such as correlation, quadrature detection, etc., may be used to separate the RF signals.

[0138] Using one or more of the techniques and features described herein, a single energized electrode on
 ⁵⁵ a staple cartridge or an isolated knife of an end-effector may be used to make multiple tissue compression measurements simultaneously. If two or more RF signals are overlaid or multiplexed (or nested or modulated), they

may be transmitted down a single power side of the endeffector and may return on either the channel frame or the anvil of the end-effector. If a filter were built into the anvil and channel contacts before they join a common return path, the tissue impedance represented by both paths could be differentiated. This may provide a measure of vertical tissue vs lateral tissue compression. This approach also may provide proximal and distal tissue compression depending on placement of the filters and location of the metallic return paths. A frequency generator and signal processor may be located on one or more chips on a circuit board or a sub board (which may already exist in an endocutter).

[0139] In various aspects, the present disclosure provides techniques for monitoring the speed and precision incrementing of the drive motor in the surgical instrument 10 (described in connection with FIGS. 1-18). In one example, a magnet can be placed on a planet frame of one of the stages of gear reduction with an inductance sensor on the gear housing. In another example, placing the magnet and magnetic field sensor on the last stage would provide the most precise incremental movement monitoring.

[0140] Conventional motor control systems employ encoders to detect the location and speed of the motor in hand held battery powered endosurgical instruments such as powered endocutter/stapler devices. Precision operation of endocutter/stapler devices relies in part on the ability to verify the motor operation under load. Simple sensor implementations may be employed to achieve verify the motor operation under load.

[0141] Accordingly, the present disclosure includes a magnetic body on one of the planetary carriers of a gear reduction system or employ brushless motor technology. Both approaches involve the placement of an inductance sensor on the outside housing of the motor or planetary gear system. In the case of a brushless motor there are electromagnetic field coils (windings, inductors, etc.) arrayed radially around the center magnetic shaft of the motor. The coils are sequentially activated and deactivated to drive the central motor shaft. One or more inductance sensors can be placed outside of the motor and adjacent to at least some of the coils to sense the activation/deactivation cycles of the motor windings to determine the number times the shaft has been rotated. Alternatively, a permanent magnet can be placed on one of the planetary carriers and the inductance sensor can be placed adjacent to the radial path of the planetary carrier to measure the number of times that stage of the gear train is rotated. This implementation can be applied to any rotational components in the system with increasingly more resolution possible in regions with a relatively large number of rotations during function, or as the rotational components become closer (in terms of number of connections) to the end effector depending on the design. The gear train sensing method may be preferred since it actually measures rotation of one of the stages whereas the motor sensing method senses the number of times

the motor has been commanded to energize, rather than the actual shaft rotation. For example, if the motor is stalled under high load, the motor sensing method would not be able to detect the lack of rotation because it senses

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only the energizing cycles not shaft rotation. Nevertheless, both techniques can be employed in a cost effective manner to sense motor rotation.

[0142] During stapling, for example, tissue is firmly clamped between opposing jaws before a staple is driven

10 into the clamped tissue. Tissue compression during clamping can cause fluid to be displaced from the compressed tissue, and the rate or amount of displacement varies depending on tissue type, tissue thickness, the surgical operation (e.g., clamping pressure and clamping

15 time). In various instances, fluid displacement between the opposing jaws of an end effector may contribute to malformation (e.g., bending) of staples between the opposing jaws. Accordingly, in various instances, it may be desirable to control the firing stroke, e.g., to control the 20 firing speed, in relationship to the detected fluid flow, or

lack thereof, intermediate opposing jaws of a surgical end effector.

[0143] Accordingly, also provided herein are methods, devices, and systems for monitoring speed and incre-25 mental movement of a surgical instrument drive train, which in turn provides information about the operational velocity of the device (e.g., jaw closure, stapling). In accordance with the present examples, the surgical instrument 10 (FIGS. 1-4) does not include a motor encoder. 30 Rather, the surgical instrument 10 may be equipped with

a motor comprising a speed sensor assembly for a power train of the motor, in accordance with an illustrative example. The speed sensor assembly can include a motor having an output shaft that is coupled directly or indirectly

35 to a drive shaft. In some examples, the output shaft is connected to a gear reduction assembly, such as a planetary gear train comprising a sensor that detects the rotational speed of any suitable component of the system. For example, the sensor may be a proximity sensor, such

40 as an induction sensor, which detects movement of one or more detectable elements affixed to any rotating part of the gear reduction assembly. The detectable element is affixed to the last stage annular gear and the sensor is positioned adjacent the radial path of the detectable

45 element so as to detect movement of the detectable element. Rotating components may vary depending on design-and the sensor(s) can be affixed to any rotating component of the gear reduction assembly. For example, in another example, a detectable element is associated with

50 the carrier gear of the final stage or even the drive gear. In some examples, a detectable element is located outside of the gear reduction assembly, such as on the driveshaft between gear reduction assembly and the end effector. In some example, a detectable element is located 55 on a rotating component in the final gear reduction at the end effector.

[0144] Various functions may be implemented utilizing the circuitry previously described, For example, the motor

may be controlled with a motor controller similar those described in connection with FIGS. 16A, 16B, 17A, 17B, and 18, where the encoder is replaced with the monitoring speed control and precision incrementing of motor systems for powered surgical instruments described herein. [0145] In one aspect, the present disclosure provides a surgical instrument 10 (described in connection with FIGS. 1-18) configured with various sensing systems. Accordingly, for conciseness and clarity the details of operation and construction will not be repeated here. In one aspect, the sensing system includes a viscoelasticity/rate of change sensing system to monitor knife acceleration, rate of change of impedance, and rate of change of tissue contact. In one example, the rate of change of knife acceleration can be used as a measure of for tissue type. In another example, the rate of change of impedance can be measures with a pulse sensor ad can be employed as a measure for compressibility. Finally, the rate of change of tissue contact can be measured with a sensor based on knife firing rate to measure tissue flow.

[0146] The rate of change of a sensed parameter or stated otherwise, how much time is necessary for a tissue parameter to reach an asymptotic steady state value, is a separate measurement in itself and may be more valuable than the sensed parameter it was derived from. To enhance measurement of tissue parameters such as waiting a predetermined amount of time before making a measurement, the present disclosure provides a novel technique for employing the derivate of the measure such as the rate of change of the tissue parameter.

[0147] The derivative technique or rate of change measure becomes most useful with the understanding that there is no single measurement that can be employed alone to dramatically improve staple formation. It is the combination of multiple measurements that make the measurements valid. In the case of tissue gap it is helpful to know how much of the jaw is covered with tissue to make the gap measure relevant. Rate of change measures of impedance may be combined with strain measurements in the anvil to relate force and compression applied to the tissue grasped between the jaw members of the end effector such as the anvil and the staple cartridge. The rate of change measure can be employed by the endosurgical device to determine the tissue type and not merely the tissue compression. Although stomach and lung tissue sometimes have similar thicknesses, and even similar compressive properties when the lung tissue is calcified, an instrument may be able to distinguish these tissue types by employing a combination of measurements such as gap, compression, force applied, tissue contact area, and rate of change of compression or rate of change of gap. If any of these measurements were used alone, the endosurgical it may be difficult for the endosurgical device to distinguish one tissue type form another. Rate of change of compression also may be helpful to enable the device to determine if the tissue is "normal" or if some abnormality exists. Measuring not only how much time has passed but the variation of the

sensor signals and determining the derivative of the signal would provide another measurement to enable the endosurgical device to measure the signal. Rate of change information also may be employed in determining when a steady state has been achieved to signal the next step in a process. For example, after clamping the tissue between the jaw members of the end effector such as the anvil and the staple cartridge, when tissue compression reaches a steady state (e.g., about 15 seconds), an indicator or trigger to start firing the device can be ena-

10 bled.

[0148] Also provided herein are methods, devices, and systems for time dependent evaluation of sensor data to determine stability, creep, and viscoelastic characteris-

15 tics of tissue during surgical instrument operation. A surgical instrument 10, such as the stapler illustrated in FIG. 1, can include a variety of sensors for measuring operational parameters, such as jaw gap size or distance, firing current, tissue compression, the amount of the jaw that

20 is covered by tissue, anvil strain, and trigger force, to name a few. These sensed measurements are important for automatic control of the surgical instrument and for providing feedback to the clinician.

[0149] The examples shown in connection with FIGS. 25 22A-37 may be employed to measure the various derived parameters such as gap distance versus time, tissue compression versus time, and anvil strain versus time. Motor current may be monitored employing the current sensor 2312 in series with the battery 2308 as described 30 herein, the current sensor 2412 in series with the battery

2408 or the current sensor 3027 in FIG. 18. [0150] FIG. 38 illustrates a motor-driven surgical instrument 8010 for cutting and fastening that may or may not be reused. The surgical instrument 8010 is similarly constructed and equipped as the surgical instrument 10 for cutting and fastening described in connection with FIGS. 1-18. In the example illustrated in FIG. 38, the surgical instrument 8010 includes a housing 8012 that comprises a handle assembly 8014 that is configured to be 40 grasped, manipulated and actuated by the clinician. The

housing 8012 is configured for operable attachment to an interchangeable shaft assembly 8200 that has an end effector 8300 operably coupled thereto that is configured to perform one or more surgical tasks or procedures.

45 Since the surgical instrument 8010 is similarly constructed and equipped as the surgical instrument 10 for cutting and fastening described in connection with FIGS. 1-18, for conciseness and clarity the details of operation and construction will not be repeated here.

50 [0151] The housing 8012 depicted in FIG. 38 is shown in connection with an interchangeable shaft assembly 8200 that includes an end effector 8300 that comprises a surgical cutting and fastening device that is configured to operably support a surgical staple cartridge 8304 55 therein. The housing 8012 may be configured for use in connection with interchangeable shaft assemblies that include end effectors that are adapted to support different sizes and types of staple cartridges, have different shaft

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lengths, sizes, and types, etc. In addition, the housing 8012 also may be effectively employed with a variety of other interchangeable shaft assemblies including those assemblies that are configured to apply other motions and forms of energy such as, for example, radio frequency (RF) energy, ultrasonic energy and/or motion to end effector arrangements adapted for use in connection with various surgical applications and procedures. Furthermore, the end effectors, shaft assemblies, handles, surgical instruments, and/or surgical instrument systems can utilize any suitable fastener, or fasteners, to fasten tissue. For instance, a fastener cartridge comprising a plurality of fasteners removably stored therein can be removably inserted into and/or attached to the end effector of a shaft assembly.

[0152] Turning now to FIG. 38, the surgical instrument 8010 is depicted that may or may not be reused. The surgical instrument 8010 is similarly constructed and equipped as the surgical instrument 10 for cutting and fastening described herein. In the example illustrated in FIG. 38, the surgical instrument 8010 includes a housing 8012 that comprises a handle assembly 8014 that is configured to be grasped, manipulated and actuated by the clinician. The housing 8012 is configured for operable attachment to an interchangeable shaft assembly 8200 that has an end effector 8300 operably coupled thereto that is configured to perform one or more surgical tasks or procedures. Since the surgical instrument 8010 is similarly constructed and equipped as the surgical instrument 10 for cutting and fastening described herein in connection with FIGS. 1-18, for conciseness and clarity the details of operation and construction will not be repeated here.

[0153] The housing 8012 depicted in FIG. 38 is shown in connection with an interchangeable shaft assembly 8200 that includes an end effector 8300 that comprises a surgical cutting and fastening device that is configured to operably support a surgical staple cartridge 8304 therein. The housing 8012 may be configured for use in connection with interchangeable shaft assemblies that include end effectors that are adapted to support different sizes and types of staple cartridges, have different shaft lengths, sizes, and types, etc. In addition, the housing 8012 also may be effectively employed with a variety of other interchangeable shaft assemblies including those assemblies that are configured to apply other motions and forms of energy such as, for example, radio frequency (RF) energy, ultrasonic energy and/or motion to end effector arrangements adapted for use in connection with various surgical applications and procedures. Furthermore, the end effectors, shaft assemblies, handles, surgical instruments, and/or surgical instrument systems can utilize any suitable fastener, or fasteners, to fasten tissue. For instance, a fastener cartridge comprising a plurality of fasteners removably stored therein can be removably inserted into and/or attached to the end effector of a shaft assembly.

[0154] FIG. 38 illustrates the surgical instrument 8010

with an interchangeable shaft assembly 8200 operably coupled thereto. In the illustrated arrangement, the handle housing forms a pistol grip portion 8019 that can be gripped and manipulated by the clinician. The handle as-

⁵ sembly 8014 operably supports a plurality of drive systems therein that are configured to generate and apply various control motions to corresponding portions of the interchangeable shaft assembly that is operably attached thereto. Trigger 8032 is operably associated with the pistol grip for controlling various of these control motions.

 tol grip for controlling various of these control motions.
 [0155] With continued reference to FIG. 38, the interchangeable shaft assembly 8200 includes an end effector 8300 that comprises an elongated channel 8302 that is configured to operably support a surgical staple car tridge 8304 therein. The end effector 8300 may further

tridge 8304 therein. The end effector 8300 may further include an anvil 8306 that is pivotally supported relative to the elongated channel 8302.

[0156] The inventors have discovered that derived parameters can be even more useful for controlling a sur-20 gical instrument, such as the instrument illustrated in FIG. 38, than the sensed parameter(s) upon which the derived parameter is based. Non-limiting examples of derived parameters include the rate of change of a sensed parameter (e.g., jaw gap distance) and how much time 25 elapses before a tissue parameter reaches an asymptotic steady state value (e.g., 15 seconds). Derived parameters, such as rate of change, are particularly useful because they dramatically improve measurement accuracy and also provide information not otherwise evident di-30 rectly from sensed parameters. For example, impedance (i.e., tissue compression) rate of change can be combined with strain in the anvil to relate compression and force, which enables the controller to determine the tissue type and not merely the amount of tissue compres-

³⁵ sion. This example is illustrative only, and any derived parameters can be combined with one or more sensed parameters to provide more accurate information about tissue types (e.g., stomach vs. lung), tissue health (calcified vs. normal), and operational status of the surgical

40 device (e.g., clamping complete). Different tissues have unique viscoelastic properties and unique rates of change, making these and other parameters discussed herein useful indicia for monitoring and automatically adjusting a surgical procedure.

45 [0157] Specifically, referring to FIGS. 38 and 39, the gap 8040 is the distance between the anvil 8306 and the elongated channel 8302 of the end effector 8300. In the open jaw position, at time zero, the gap 8040 between the anvil 8306 and the elongated member is at its max-50 imum distance. The width of the gap 8040 decreases as the anvil 8306 closes, such as during tissue clamping. The gap distance rate of change can vary because tissue has non-uniform resiliency. For example, certain tissue types may initially show rapid compression, resulting in 55 a faster rate of change. However, as tissue is continually compressed, the viscoelastic properties of the tissue can cause the rate of change to decrease until the tissue cannot be compressed further, at which point the gap dis-

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tance will remain substantially constant. The gap decreases over time as the tissue is squeezed between the anvil 8306 and the surgical staple cartridge 8304 of the end effector 8300. The one or more sensors described in connection with FIGS. 22A-37 and FIG. 40 may be adapted and configured to measure the gap distance "d" between the anvil 8306 and the surgical staple cartridge 8304 over time t and the rate of change of the gap distance "d" over time t is the Slope of the curve, where Slope = $\Delta d / \Delta t$. In addition, the rate of change of firing current is can be used as an indicator that the tissue is transitioning from one state to another state. Accordingly, firing current and, in particular, the rate of change of firing current can be used to monitor device operation. The firing current decreases over time as the knife cuts through the tissue. The rate of change of firing current can vary if the tissue being cut provides more or less resistance due to tissue properties or sharpness of the knife 8305 (FIG. 39). For example, the motor current may be monitored employing the current sensor 2312 in series with the battery 2308 as described herein, the current sensor 2412 in series with the battery 2408 shown herein, or the current sensor 3027 shown in FIG. 18. The current sensors 2312, 2314, 3027 may be adapted and configured to measure the motor firing current "i" over time t and the rate of change of the firing current "i" over time t is the Slope of the curve, where Slope = $\Delta i / \Delta t$. The sensors described in connection with FIGS. 22A-37 and 40 may be adapted and configured to measure tissue compression/impedance. The sensors may be adapted and configured to measure tissue impedance "Z" over time t and the rate of change of the tissue impedance "Z" over time t is the Slope, where Slope = $\Delta Z / \Delta t$. The rate of change of anvil 8306 strain can be measured by a pressure sensor or strain gauge positioned on either or both the anvil 8306 and the surgical staple cartridge 8304 (FIGS. 38, 39) to measure the pressure or strain applied to the tissue grasped between the anvil 8306 and the surgical staple cartridge 8304. Thus, at time zero, trigger 8020 (FIG. 38) pressure may be at its lowest and trigger pressure may increase until completion of an operation (e.g., clamping, cutting, or stapling). The rate of change trigger force can be measured by a pressure sensor or strain gauge positioned on the trigger 8032 of the pistol grip portion 8019 of the handle of the surgical instrument 8010 (FIG. 38) to measure the force required to drive the knife 8305 (FIG. 39) through the tissue grasped between the anvil 8306 and the surgical staple cartridge 8304.

[0158] Turning briefly to FIG. 40, the end effector 9012 is one aspect of the end effector 8300 (FIG. 38) that may be adapted to operate with surgical instrument 8010 (FIG. 38) to measure the various derived parameters such as gap distance versus time, tissue compression versus time, and anvil strain versus time. Accordingly, the end effector 9012 shown in FIG. 40 may include one or more sensors configured to measure one or more parameters or characteristics associated with the end effector 9012 and/or a tissue section captured by the end

effector 9012. In the example illustrated in FIG. 40, the end effector 9012 comprises a first sensor 9020 and a second sensor 9026. In various examples, the first sensor 9020 and/or the second sensor 9026 may comprise, for example, a magnetic sensor such as, for example, a magnetic field sensor, a strain gauge, a pressure sensor, a force sensor, an inductive sensor such as, for example, an eddy current sensor, a resistive sensor, a capacitive sensor, an optical sensor, and/or any other suitable sensor for measuring one or more parameters of the end

effector 9012. **[0159]** In certain instances, the first sensor 9020 and/or the second sensor 9026 may comprise, for example, a magnetic field sensor embedded in an anvil 9014 and

¹⁵ configured to detect a magnetic field generated by a magnet 9024 embedded in a jaw member 9016 and/or the staple cartridge 9018. The anvil 9014 is pivotally rotatable between open and closed positions. The strength of the detected magnetic field may correspond to, for example,

the thickness and/or fullness of a bite of tissue located between the anvil 9014 and the jaw member 9016. In certain instances, the first sensor 9020 and/or the second sensor 9026 may comprise a strain gauge, such as, for example, a micro-strain gauge, configured to measure the magnitude of the strain in the anvil 9014 during a clamped condition. The strain gauge provides an electri-

clamped condition. The strain gauge provides an electrical signal whose amplitude varies with the magnitude of the strain.
 [0160] In some aspects, one or more sensors of the

30 end effector 9012 such as, for example, the first sensor 9020 and/or the second sensor 9026 may comprise a pressure sensor configured to detect a pressure generated by the presence of compressed tissue between the anvil 9014 and the jaw member 9016. In some examples,

one or more sensors of the end effector 9012 such as, for example, the first sensor 9020 and/or the second sensor 9026 are configured to detect the impedance of a tissue section located between the anvil 9014 and the jaw member 9016. The detected impedance may be indicative of the thickness and/or fullness of tissue located between the anvil 9014 and the jaw member 9016.

[0161] In one aspect, one or more of the sensors of the end effector 9012 such as, for example, the first sensor 9020 is configured to measure the gap 9022 between

45 the anvil 9014 and the jaw member 9016. In certain instances, the gap 9022 can be representative of the thickness and/or compressibility of a tissue section clamped between the anvil 9014 and the jaw member 9016. In at least one example, the gap 9022 can be equal, or sub-50 stantially equal, to the thickness of the tissue section clamped between the anvil 9014 and the jaw member 9016. In one example, one or more of the sensors of the end effector 9012 such as, for example, the first sensor 9020 is configured to measure one or more forces exert-55 ed on the anvil 9014 by the jaw member 9016 and/or tissue clamped between the anvil 9014 and the jaw member 9016. The forces exerted on the anvil 9014 can be representative of the tissue compression experienced by

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the tissue section captured between the anvil 9014 and the jaw member 9016. In one aspect, the gap 9022 between the anvil 9014 and the jaw member 9016 can be measured by positioning a magnetic field sensor on the anvil 9014 and positioning a magnet on the jaw member 9016 such that the gap 9022 is proportional to the signal detected by the magnetic field sensor and the signal is proportional to the distance between the magnet and the magnetic field sensor. It will be appreciated that the location of the magnetic field sensor and the magnet may be swapped such that the magnetic field sensor is positioned on the jaw member 9016 and the magnet is placed on the anvil 9014.

[0162] One or more of the sensors such as, for example, the first sensor 9020 and/or the second sensor 9026 may be measured in real-time during a clamping operation. Real -time measurement allows time based information to be analyzed, for example, by a processor, and used to select one or more algorithms and/or look-up tables for the purpose of assessing, in real -time, a manual input of an operator of the surgical instrument 9010. Furthermore, real -time feedback can be provided to the operator to assist the operator in calibrating the manual input to yield a desired output.

[0163] FIG. 41 is a logic diagram illustrating one aspect of a real-time feedback system 9060 for assessing, in real -time, a manual input 9064 of an operator of the surgical instrument 9010 and providing to the operator real -time feedback as to the adequacy of the manual input 9064. With reference to FIGS. 40 and 41, in the example illustrated in FIG. 41, the real-time feedback system 9060 is comprised of a circuit. The circuit includes a controller 9061 comprising a processor 9062. A sensor such as, for example, the first sensor 9020 is employed by the processor 9062 to measure a parameter of the end effector 9012. In addition, the processor 9062 can be configured to determine or receive a value representative of a manual input 9064 of an operator of the surgical instrument 9010. The manual input 9064 can be continuously assessed by the processor 9062 for as long as the manual input 9064 is being provided by the operator. The processor 9062 can be configured to monitor a value representative of the manual input 9064. Furthermore, the processor 9062 is configured to assign, select, or determine a position, rank, and/or status for the determined value with respect to a desired zone or range. The measurement of the parameter of the end effector 9012 and the determined value can be employed by the processor 9062 to select or determine the position, rank, and/or status associated with the determined value, as described in greater detail below. A change in the manual input 9064 yields a change in the determined value which, in turn, yields a change in the position, rank, and/or status assigned to the determined value with respect to the desired zone or range.

[0164] As illustrated in FIG. 41, the real-time feedback system 9060 may further include a feedback indicator 9066 which can be adjusted between a plurality of posi-

tions, ranks, and/or statuses inside and outside a desired zone or range. In one example, the processor 9062 may select a first position (P1), rank, and/or status that characterizes the manual input 9064 based on a measurement (M1) of a parameter of the end effector 9012 and a first determined value (VI) representing a first manual input (I1). In certain instances, the first position (P1), rank, and/or status may fall outside the desired zone or range. In such instances, the operator may change the manual

¹⁰ input 9064 from the first manual input (I1) to a second manual input (I2) by increasing or decreasing the manual input 9064, for example. In response, the processor 9062 may adjust the feedback indicator 9066 from the first position (P1), rank, and/or status to a second position (P2),

rank, and/or status, which characterizes the change to the manual input 9064. The processor 9062 may select the second position (P2), rank, and/or status based on the measurement (M1) of the parameter of the end effector 9012 and a second determined value (V2) representing a second manual input (I2). In certain instances, the second position (P2), rank, and/or status may fall inside the desired zone or range. In such instances, the operator may maintain the second manual input (I2) for a remainder of a treatment cycle or procedure, for example.

ple.
[0165] In the aspect illustrated in FIG. 41, the controller 9061 includes a storage medium such as, for example, a memory 9068. The memory 9068 may be configured to store correlations between measurements of one or more parameters of the end effector 9012, values representing manual inputs, and corresponding positions, ranks, and/or statuses characterizing the manual input 9064 with respect to a desired zone or range. In one example, the memory 9068 may store the correlation between the measurement (M1), the first determined value (V1), and the first manual input (I1), and the correlation

between the measurement (M1), the second determined value (V2), and the second manual input (I2). In one example, the memory 9068 may store an algorism, an equa-

40 tion, or a look-up table for determining correlations between measurements of one or more parameters of the end effector 9012, values representing manual inputs, and corresponding positions, ranks, or statuses with respect to a desired zone or range. The processor 9062

⁴⁵ may employ such algorism, equation, and/or look-up table to characterize a manual input 9064 provided by an operator of the surgical instrument 9010 and provide feedback to the operator as to the adequacy of the manual input 9064.

 ⁵⁰ [0166] FIG. 42 is a logic diagram illustrating one aspect of a real-time feedback system 9070. The real-time feedback system 9070 is similar in many respects to the realtime feedback system 9060. For example, like the realtime feedback system 9060, the real-time feedback system 9070 is configured for assessing, in real-time, a manual input of an operator of the surgical instrument 9010 and providing to the operator real -time feedback as to the adequacy of the manual input. Furthermore, like the

[0167] In the aspect illustrated in FIG. 42, a sensor 9072, such as, for example, a strain gauge or a microstrain gauge, is configured to measure one or more parameters of the end effector 9012, such as, for example, the amplitude of the strain exerted on the anvil 9014 during a clamping operation, which can be indicative of the tissue compression. The measured strain is converted to a digital signal and provided to the processor 9062. A sensor 9074, such as, for example, a load sensor, can measure the force to advance the cutting member 9040 to cut tissue captured between the anvil 9014 and the staple cartridge 9018. Alternatively, a current sensor (not shown) can be employed to measure the current drawn by the motor 9082. The force required to advance the firing bar 9036 can correspond to the current drawn by the motor 9082, for example. The measured force is converted to a digital signal and provided to the processor 9062. A sensor 9076, such as, for example, a magnetic field sensor, can be employed to measure the thickness of the captured tissue, as described above. The measurement of the magnetic field sensor 9076 is also converted to a digital signal and provided to the processor 9062.

[0168] In the aspect illustrated in FIG. 42, the real-time feedback system 9070 further includes the tracking system 9080 which can be configured to determine the position of the firing trigger. As described above, the firing trigger 9094 can be depressed or actuated by moving the firing trigger 9094 between a plurality of positions, each corresponding to one of a plurality of values of a characteristic of motion of the firing bar 9036 and/or the cutting member 9040 during a firing stroke. As describe above, a characteristic of motion can be a speed of advancement of the firing bar 9036 and/or the cutting member 9040 during the firing stroke. In certain instances, a motor driver 9092 can be in communication with the controller 9061, and can be configured to drive the motor 9082 in accordance with an operator's manual input as detected by the tracking system 9080.

[0169] Further to the above, the real-time feedback system 9070 may include a feedback indicator 9066. In one aspect, the feedback indicator 9066 can be disposed in the handle 9030. Alternatively, the feedback indicator can be disposed in the shaft assembly 9032, for example. In any event, the controller 9061 may employ the feedback indicator 9066 to provide feedback to an operator of the surgical instrument 9010 with regard to the adequacy of a manual input such as, for example, a selected position of the firing trigger 9094. To do so, the controller 9061 may assess the selected position of the firing trigger 9094 and/or the corresponding value of the speed of the firing bar 9036 and/or the cutting member 9040. The measurements of the tissue compression, the tissue thickness, and/or the force required to advance the firing bar 9036, as respectively measured by the sensors 9072, 9074, and 9076, can be used by the controller 9061 to characterize the selected position of the firing trigger 9094 and/or the corresponding value of the speed of the firing bar 9036 and/or the cutting member 9040. In one instance, the memory 9068 may store an algorism, an equation, and/or a look-up table which can be employed by the controller 9061 in the assessment. In one example, the measurements of the sensors 9072, 9074, and/or 9076 can be used to select or determine a position, rank,

- ¹⁰ and/or a status that characterizes the selected position of the firing trigger 9094 and/or the corresponding value of the speed of the firing bar 9036 and/or the cutting member 9040. The determined position, rank, and/or status can be communicated to the operator via the feedback ¹⁵ indicator 9066.
- [0170] The reader will appreciate that an optimal speed of the firing bar 9036 and/or the cutting member 9040 during a firing stroke can depend on several parameters of the end effector 9012 such as, for example, the thick²⁰ ness of the tissue captured by the end effector 9012, the tissue compression, and/or the force required to advance the firing bar 9036 and, in turn, the cutting member 9040. As such, measurements of these parameters can be leveraged by the controller 9061 in assessing whether a
- ²⁵ current speed of advancement of the cutting member 9040 through the captured tissue is within an optimal zone or range.

[0171] In one aspect, a plurality of smart sensors may be positioned on a power line of an end-effector and may
³⁰ be communicatively coupled to a handle of an endocutter. The smart sensors may be positioned in series or parallel with respect to the power line. Referring now to FIG. 43, smart sensors 12060 and 12062 may be in communication with a signal processing component or a proc³⁵ essor 12064 which may be local to the smart sensors.

- Both the smart sensors 12060 and 12062 and the processor 12064 may be located at the end-effector (represented by dashed-box 12066). For example, smart sensor 12060 may output signals or data to an operational amplifier 12068 and an ADC converter 12070, which may
- 40 amplifier 12068 and an ADC converter 12070, which may condition the signals or data for input into processor 12064. Similarly, smart sensor 12062 may output signals or data to an operational amplifier 12072 and an ADC converter 12074, which may condition the signals or data
 45 for input into processor 12064.

[0172] Smart sensors 12060 and/or 12062 may be different types of sensors or the same type of sensor, which may be, for example, magnetic field sensors, magnetic sensors, inductive sensors, capacitive sensors, or other types of sensors used in medical devices or endocutters.

⁵⁰ types of sensors used in medical devices or endocutters. Component 12064, previously referred to as a processor, also may be a computational core, FPGA (field programmable gate array), logic unit (e.g., logic processor or logic controller), signal processing unit, or other type of proc-55 essor. The processor 12064 may be in communication with a memory, such as non-volatile memory 12076, which may store calculation data, equipment information such as a type of cartridge inserted in the end-effector 12066, tabular data, or other reference data that may enable the processor 12064 to process signals or data received from one or more of the smart sensors 12060 or 12062 for use in operating the end-effector 12066 or an endocutter.

[0173] Further, a shaft 12078 may include a return path through which at least one of the plurality of smart sensors (e.g., smart sensors 12060 or 12062) and the handle 12080 are communicatively coupled. The shaft may include one or more wires which may transfer information from the processor 12064 to the handle 12080 for operation of the end-effector 12066 or endocutter. In one example, the information from the processor 12064 may be communicated to the handle 12080 (by way of shaft 12078 or directly without use of shaft 12078) over one or more of: a wired-line, a single-wired line, a multi-wired line, a wireless communication protocol such as Bluetooth, an optical line, or an acoustic line.

[0174] In one aspect, at least one of a plurality of smart sensors positioned at an end-effector may include a signal processing component. For example, the signal processing component may be built into the smart sensor or may be locally coupled to the smart sensor as a single module. The signal processing component may be configured to process data received from a sensor component (e.g., sensor component 12020) of at least one of the plurality of smart sensors. A controller 12024 (e.g., a controller) at the handle may be communicatively coupled to at least one of the plurality of smart sensors.

[0175] In one aspect, a smart sensor may be configured for local signal processing in a medical device. The smart sensor may include at least one sensor component (e.g., sensor component 12020) and at least one processing component (e.g., processing component 12022). The processing component may be configured to receive data from the at least one sensor component and to process the data into information for use by the medical device. The medical device may be, for example, an endocutter, however this is not intended to be a limitation of the present disclosure. It should be understood that the techniques and features discussed herein for smart sensors with local signal processing may be used in any medical device where processing of sensor signals or data is used for operation of the medical device.

[0176] Further, a controller (e.g., controller 12024, controller) in the medical device may be configured to receive the information (i.e., processed signals or data) from the at least one processing component (e.g., processing component 12022). As discussed above, the medical device may be a surgical instrument such as an endocutter and the smart sensor may be configured for local signal processing may refer to, for example, processing signals or data from a sensor component at a processing component coupled to the sensor, where the resulting processed information may be used by a separate component. For example, the controller 12024 may be positioned in the handle 12012 of the surgical instrument (i.e.,

the endocutter 12010) and the smart sensor may be configured to be positioned in a separate component (i.e., the end-effector 12016) of the surgical instrument (i.e., the endocutter 12010), separate from the handle 12012.

⁵ Thus, the controller 12024 may be positioned at the handle 12012 of the surgical instrument and the signal processing component 12022 and the sensor 12020 may be located in a component separate from the handle 12012 (e.g., end-effector 12016).

10 [0177] In this way, the handle or controller 12024 need not have information about the smart sensor, knowledge of what the smart sensor is doing, or capability to interpret data feed back from the smart sensor. This is because the processing component 12022 may transform or con-

¹⁵ dition the data from the smart sensor and generate information from the data directly usable by the handle or controller 12024. The information generated by the processing component may be used directly, without the data from the smart sensor needing to be processed in

²⁰ another part of the medical device (e.g., near the handle 12012 or controller 12024). Thus, the surgical instrument may be controlled based on the (processed) information from the signal processing component local to the sensor.

²⁵ [0178] In one aspect, a current draw on a power line communicatively coupled to the signal processing component 12022 (i.e., local to the sensor 12020) may be monitored. The current draw may be monitored by a processor, controller, or other monitoring device at the shaft

³⁰ 12014 or the handle 12012, or at another processor, controller or other monitoring device separate from the signal processing component 12022. For example, the monitoring may be a standard Morse Code type monitoring of the current draw on the power line. An issue with the

³⁵ surgical instrument based on the current draw and a particular sensor may be determined by the separate processor at, e.g., the handle 12012. In this way, the monitoring may allow the handle (or a processor or controller therein) to be informed of various issues related to signals

40 or data received by one or more sensor and which particular sensor identified the issue, without a further communication requirement (e.g., pairing, or other coupled communication).

[0179] FIG. 44 illustrates one aspect of a circuit 13190 45 configured to convert signals from the first sensor 13158 and the plurality of secondary sensors 13160a, 13160b into digital signals receivable by a processor, such as, for example, the primary processor 2006 (FIGS. 16A-16B). The circuit 13190 comprises an analog-to-digital 50 convertor 13194. In some examples, the analog-to-digital convertor 13194 comprises a 4-channel, 18-bit analog to digital convertor. Those skilled in the art will recognize that the analog-to-digital convertor 13194 may comprise any suitable number of channels and/or bits to convert 55 one or more inputs from analog to digital signals. The circuit 13190 comprises one or more level shifting resistors 13196 configured to receive an input from the first sensor 13158, such as, for example, a magnetic field

sensor. The level shifting resistors 13196 adjust the input from the first sensor, shifting the value to a higher or lower voltage depending on the input. The level shifting resistors 13196 provide the level-shifted input from the first sensor 13158 to the analog-to-digital convertor.

[0180] In some aspects, a plurality of secondary sensors 13160a, 13160b are coupled to a plurality of bridges 13192a, 13192b within the circuit 13190. The plurality of bridges 13192a, 13192b may provide filtering of the input from the plurality of secondary sensors 13160a, 13160b. After filtering the input signals, the plurality of bridges 13192a, 13192b provide the inputs from the plurality of secondary sensors 13160a, 13160b to the analog-to-digital convertor 13194. In some examples, a switch 13198 coupled to one or more level shifting resistors may be coupled to the analog-to-digital convertor 13194. The switch 13198 is configured to calibrate one or more of the input signals, such as, for example, an input from a magnetic field sensor. The switch 13198 may be engaged to provide one or more level shifting signals to adjust the input of one or more of the sensors, such as, for example, to calibrate the input of a magnetic field sensor. In some examples, the adjustment is not necessary, and the switch 13198 is left in the open position to decouple the level shifting resistors. The switch 13198 is coupled to the analog-to-digital convertor 13194. The analog-to-digital convertor 13194 provides an output to one or more processors, such as, for example, the primary processor 2006 (FIGS. 16A-16B). The primary processor 2006 calculates one or more parameters of the end effector 13150 based on the input from the analog-to-digital convertor 13194. For example, in one example, the primary processor 2006 calculates a thickness of tissue located between the anvil 13152 and the staple cartridge 13156 based on inputs from the fist sensor 13158 and the plurality of secondary sensors 13160a, 13160b.

[0181] FIG. 45 illustrates one aspect of a staple cartridge 13606 that comprises a flex cable 13630 connected to a magnetic field sensor 13610 and processor 13612. The staple cartridge 13606 is similar to the staple cartridge 13606 is similar to the surgical staple cartridge 304 (FIG. 1) described above in connection with surgical instrument 10 (FIGS. 1-6). FIG. 45 is an exploded view of the staple cartridge 13606. The staple cartridge comprises 13606 a cartridge body 13620, a wedge sled 13618, a cartridge tray 13622, and a flex cable 13630. The flex cable 13630 further comprises electrical contacts 13632 at the proximal end of the staple cartridge 13606, placed to make an electrical connection when the staple cartridge 13606 is operatively coupled with an end effector, such as end effector 13800 described below. The electrical contacts 13632 are integrated with cable traces 13634, which extend along some of the length of the staple cartridge 13606. The cable traces 13634 connect 13636 near the distal end of the staple cartridge 13606 and this connection 13636 joins with a conductive coupling 13614. A magnetic field sensor 13610 and a processor 13612 are operatively coupled to the conductive coupling 13614 such that the magnetic field sensor 13610 and the processor 13612 are able to communicate.

- [0182] FIG. 46 illustrates one aspect of an end effector
 ⁵ 13800 with a flex cable 13830 operable to provide power to a staple cartridge 13806 that comprises a distal sensor plug 13816. The end effector 13800 is similar to the end effector 300 (FIG. 1) described above in connection with surgical instrument 10 (FIGS. 1-6). The end effector
- 13800 comprises an anvil 13802, a jaw member or elongated channel 13804, and a staple cartridge 13806 operatively coupled to the elongated channel 13804. The end effector 13800 is operatively coupled to a shaft assembly. The shaft assembly is similar to interchangeable

¹⁵ shaft assembly 200 (FIG. 1) described above in connection with surgical instrument 10 (FIGS. 1-6). The shaft assembly further comprises a closure tube that encloses the exterior of the shaft assembly. In some examples the shaft assembly further comprises an articulation joint
²⁰ 13904, which includes a double pivot closure sleeve assembly. The double pivot closure sleeve assembly includes an end effector closure sleeve assembly that is

operable to couple with the end effector 13800.

[0183] FIGS. 47 and 48 illustrate the elongated channel 13804 portion of the end effector 13800 without the anvil 13802 or the staple cartridge, to illustrate how the flex cable 13830 can be seated within the elongated channel 13804. In some examples, the elongated channel 13804 further comprises a third aperture 13824 for receiving the flex cable 13830. Within the body of the elongated channel 13804 the flex cable splits 13834 to form extensions 13836 on either side of the elongated channel 13804. FIG. 48 further illustrates that connectors 13838 can be operatively coupled to the flex cable ex-

[0184] FIG. 49 illustrates the flex cable 13830 alone. As illustrated, the flex cable 13830 comprises a single coil 13832 operative to wrap around the articulation joint 13904 (FIG. 46), and a split 13834 that attaches to extensions 13836. The extensions can be coupled to con-

40 tensions 13836. The extensions can be coupled to connectors 13838 that have on their distal facing surfaces prongs 13840 for coupling to the staple cartridge 13806, as described below.

[0185] FIG. 50 illustrates a close up view of the elon-45 gated channel 13804 shown in FIG. 47 and 48 with a staple cartridge 13804 coupled thereto. The staple cartridge 13804 comprises a cartridge body 13822 and a cartridge tray 13820. In some examples the staple cartridge 13806 further comprises electrical traces 13828 50 that are coupled to proximal contacts 13856 at the proximal end of the staple cartridge 13806. The proximal contacts 13856 can be positioned to form a conductive connection with the prongs 13840 of the connectors 13838 that are coupled to the flex cable extensions 13836. Thus, 55 when the staple cartridge 13806 is operatively coupled with the elongated channel 13804, the flex cable 13830, through the connectors 13838 and the connector prongs 13840, can provide power to the staple cartridge 13806.

[0186] FIGS. 51 and 52 illustrate one aspect of a distal sensor plug 13816. FIG. 51 illustrates a cutaway view of the distal sensor plug 13816 comprises a magnetic field sensor 13810 and a processor 13812. The distal sensor plug 13816 further comprises a flex board 13814. As further illustrated in FIG. 52, the magnetic field sensor 13810 and the processor 13812 are operatively coupled to the flex board 13814 such that they are capable of communicating.

[0187] FIG. 53 illustrates one aspect of an end effector 13950 with a flex cable 13980 operable to provide power to sensors and electronics in the distal tip 13952 of the anvil 13961 portion. The end effector 13950 comprises an anvil 13961, a jaw member or elongated channel 13954, and a staple cartridge 13956 operatively coupled to the elongated channel. The end effector 13950 is operatively coupled to a shaft assembly 13960. The shaft assembly 13960 further comprises a closure tube 13962 that encloses the shaft assembly 13960. In some examples the shaft assembly 13960 further comprises an articulation joint 13964, which includes a double pivot closure sleeve assembly 13966.

[0188] In various aspects, the end effector 13950 further comprises a flex cable 13980 that is configured to not interfere with the function of the articulation joint 13964. In some examples, the closure tube 13962 comprises a first aperture 13968 through which the flex cable 13980 can extend. In some examples, flex cable 13980 further comprises a loop or coil 13982 that wraps around the articulation joint 13964 such that the flex cable 13980 does not interfere with the operation of the articulation joint 13964, as further described below. In some examples, the flex cable 13980 extends along the length of the anvil 13961 to a second aperture 13970 in the distal tip of the anvil 13961.

[0189] A portion of a surgical stapling instrument 16000 is illustrated in FIGS. 54-56. The stapling instrument 16000 is usable with a manually-operated system and/or a robotically-controlled system, for example. The stapling instrument 16000 comprises a shaft 16010 and an end effector 16020 extending from the shaft 16010. The end effector 16020 comprises a cartridge channel 16030 and a staple cartridge 16050 positioned in the cartridge channel 16030. The staple cartridge 16050 comprises a cartridge body 16051 and a retainer 16057 attached to the cartridge body 16051. The cartridge body 16051 is comprised of a plastic material, for example, and the retainer 16057 is comprised of metal, for example; however, the cartridge body 16051 and the retainer 16057 can be comprised of any suitable material. The cartridge body 16051 comprises a deck 16052 configured to support tissue, a longitudinal slot 16056, and a plurality of staple cavities 16053 defined in the deck 16052.

[0190] Referring primarily to FIGS. 55 and 56, staples 16055 are removably positioned in the staple cavities 16053 and are supported by staple drivers 16054 which are also movably positioned in the staple cavities 16053.

The retainer 16057 extends around the bottom of the cartridge body 16051 to keep the staple drivers 16054 and/or the staples 16055 from falling out of the bottom of the staple cavities 16053. The staple drivers 16054 and the staples 16055 are movable between an unfired position (FIG. 55) and a fired position by a sled 16060.

The sled 16060 is movable between a proximal, unfired position (FIG. 55) toward a distal, fired position to eject the staples 16055 from the staple cartridge 16050, as

¹⁰ illustrated in FIG. 56. The sled 16060 comprises one or more ramped surfaces 16064 which are configured to slide under the staple drivers 16054. The end effector 16020 further comprises an anvil 16040 configured to deform the staples 16055 when the staples 16055 are

¹⁵ ejected from the staple cartridge 16050. In various instances, the anvil 16040 can comprise forming pockets 16045 defined therein which are configured to deform the staples 16055.

[0191] The shaft 16010 comprises a frame 16012 and an outer sleeve 16014 which is movable relative to the frame 16012. The cartridge channel 16030 is mounted to and extends from the shaft frame 16012. The outer sleeve 16014 is operably engaged with the anvil 16040 and is configured to move the anvil 16040 between an

open position (FIG. 54) and a closed position (FIG. 55).
 In use, the anvil 16040 is movable toward a staple cartridge 16050 positioned in the cartridge channel 16030 to clamp tissue against the deck 16052 of the staple cartridge 16050. In various alternative aspects, the cartridge
 channel 16030 and the staple cartridge 16050 are mov-

able relative to the anvil 16040 to clamp tissue therebetween. In either event, the shaft 16010 further comprises a firing member 16070 configured to push the sled 16060 distally. The firing member 16070 comprises a knife edge

³⁵ 16076 which is movable within the longitudinal slot 16056 and is configured to incise the tissue positioned intermediate the anvil 16040 and the staple cartridge 16050 as the firing member 16070 is advanced distally to eject the staples 16055 from the staple cartridge 16050. The firing
⁴⁰ member 16070 further comprises a first cam 16071 configured to engage the cartridge channel 16030 and a second cam 16079 configured to engage the anvil 16040

and hold the anvil 16040 in position relative to the staple cartridge 16050. The first cam 16071 is configured to
⁴⁵ slide under the cartridge channel 16030 and the second cam 16079 is configured to slide within an elongated slot 16049 defined in the anvil 16040.

[0192] FIG. 57 illustrates one aspect of an end effector 3011 comprising a first sensor 3008a and a second sensor 3008b. The end effector 3011 is similar to the end effector 300 described above. The end effector 3011 comprises an anvil 3013 pivotally coupled to a jaw member 3004. The jaw member 3004 is configured to receive a staple cartridge 3021 therein. The staple cartridge 3021
⁵⁵ comprises a plurality of staples (not shown). The plurality of staples is deployable from the staple cartridge 3021 during a surgical operation. The end effector 3011 comprises a first sensor 3008a configured to measure one

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or more parameters of the end effector 3011. For example, in one aspect, the first sensor 3008a is configured to measure the gap 3023 between the anvil 3013 and the jaw member 3004. The first sensor 3008a may comprise, for example, a Hall effect sensor configured to detect a magnetic field generated by a magnet 3012 embedded in the second jaw member 3004 and/or the staple cartridge 3021. As another example, in one aspect, the first sensor 3008a is configured to measure one or more forces exerted on the anvil 3013 by the second jaw member 3004 and/or tissue clamped between the anvil 3013 and the second jaw member 3004.

[0193] The end effector 3011 comprises a second sensor 3008b. The second sensor 3008b is configured to measure one or more parameters of the end effector 3011. For example, in various aspects, the second sensor 3008b may comprise a strain gauge configured to measure the magnitude of the strain in the anvil 3013 during a clamped condition. The strain gauge provides an electrical signal whose amplitude varies with the magnitude of the strain. In various aspects, the first sensor 3008a and/or the second sensor 3008b may comprise, for example, a magnetic sensor such as, for example, a Hall effect sensor, a strain gauge, a pressure sensor, a force sensor, an inductive sensor such as, for example, an eddy current sensor, a resistive sensor, a capacitive sensor, an optical sensor, and/or any other suitable sensor for measuring one or more parameters of the end effector 3011. The first sensor 3008a and the second sensor 3008b may be arranged in a series configuration and/or a parallel configuration. In a series configuration, the second sensor 3008b may be configured to directly affect the output of the first sensor 3008a. In a parallel configuration, the second sensor 3008b may be configured to indirectly affect the output of the first sensor 3008a.

[0194] In one aspect, the one or more parameters measured by the first sensor 3008a are related to the one or more parameters measured by the second sensor 3008b. For example, in one aspect, the first sensor 3008a is configured to measure the gap 3023 between the anvil 3013 and the jaw member 3004. The gap 3023 is representative of the thickness and/or compressibility of a tissue section clamped between the anvil 3013 and the staple cartridge 3021 located in the jaw member 3004. The first sensor 3008a may comprise, for example, a Hall effect sensor configured to detect a magnetic field generated by a magnet 3012 coupled to the second jaw member 3004 and/or the staple cartridge 3021. Measuring at a single location accurately describes the compressed tissue thickness for a calibrated full bit of tissue, but may provide inaccurate results when a partial bite of tissue is placed between the anvil 3013 and the second jaw member 3004. A partial bite of tissue, either a proximal partial bite or a distal partial bite, changes the clamping geometry of the anvil 3013.

[0195] In some aspects, the second sensor 3008b is configured to detect one or more parameters indicative

of a type of tissue bite, for example, a full bite, a partial proximal bite, and/or a partial distal bite. The measurement of the second sensor 3008b may be used to adjust the measurement of the first sensor 3008a to accurately represent a proximal or distal positioned partial bite's true compressed tissue thickness. For example, in one aspect, the second sensor 3008b comprises a strain gauge,

such as, for example, a micro-strain gauge, configured to monitor the amplitude of the strain in the anvil during a clamped condition. The amplitude of the strain of the

anvil 3013 is used to modify the output of the first sensor 3008a, for example, a Hall effect sensor, to accurately represent a proximal or distal positioned partial bite's true compressed tissue thickness. The first sensor 3008a and

¹⁵ the second sensor 3008b may be measured in real-time during a clamping operation. Real-time measurement allows time based information to be analyzed, for example, by the primary processor 2006, and used to select one or more algorithms and/or look-up tables to recognize
²⁰ tissue characteristics and clamping positioning to dy-

namically adjust tissue thickness measurements. [0196] In some aspects, the thickness measurement of the first sensor 3008a may be provided to an output device of a surgical instrument 10 coupled to the end 25 effector 3011. For example, in one aspect, the end effector 3011 is coupled to the surgical instrument 10 comprising a display 2028. The measurement of the first sensor 3008a is provided to a processor, for example, the primary processor 2006. The primary processor 2006 ad-30 justs the measurement of the first sensor 3008a based on the measurement of the second sensor 3008b to reflect the true tissue thickness of a tissue section clamped between the anvil 3013 and the staple cartridge 3021. The primary processor 2006 outputs the adjusted tissue 35 thickness measurement and an indication of full or partial bite to the display 2028. An operator may determine whether or not to deploy the staples in the staple cartridge 3021 based on the displayed values.

[0197] In some aspects, the first sensor 3008a and the
 second sensor 3008b may be located in different environments, such as, for example, the first sensor 3008a being located within a patient at a treatment site and the second sensor 3008b being located externally to the patient. The second sensor 3008b may be configured to

⁴⁵ calibrate and/or modify the output of the first sensor 3008a. The first sensor 3008a and/or the second sensor 3008b may comprise, for example, an environmental sensor. Environmental sensors may comprise, for example, temperature sensors, humidity sensors, pressure sensors, and/or any other suitable environmental sensor.

[0198] FIG. 58 is a logic diagram illustrating one aspect of a process 3050 for determining and displaying the thickness of a tissue section clamped between the anvil 3013 and the staple cartridge 3021 of the end effector 3011. The process 3050 comprises obtaining a Hall effect voltage 3052, for example, through a Hall effect sensor located at the distal tip of the anvil 3013. The Hall effect voltage 3052 is provided to an analog to digital convertor

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3054 and converted into a digital signal. The digital signal is provided to a processor, such as, for example, the primary processor 2006. The primary processor 2006 calibrates 3056 the curve input of the Hall effect voltage 3052 signal. A strain gauge 3058, such as, for example, a micro-strain gauge, is configured to measure one or more parameters of the end effector 3011, such as, for example, the amplitude of the strain exerted on the anvil 3013 during a clamping operation. The measured strain is converted 3060 to a digital signal and provided to the processor, such as, for example, the primary processor 2006. The primary processor 2006 uses one or more algorithms and/or lookup tables to adjust the Hall effect voltage 3052 in response to the strain measured by the strain gauge 3058 to reflect the true thickness and fullness of the bite of tissue clamped by the anvil 3013 and the staple cartridge 3021. The adjusted thickness is displayed 3026 to an operator by, for example, a display 2026 embedded in the surgical instrument 10.

[0199] In some aspects, the surgical instrument can further comprise a load sensor 3082 or load cell. The load sensor 3082 can be located, for instance, in the interchangeable shaft assembly 200, described above, or in the housing 12, also described above.

[0200] FIG. 59 is a logic diagram illustrating one aspect of a process 3070 for determining and displaying the thickness of a tissue section clamped between the anvil 3013 and the staple cartridge 3021 of the end effector 3011. The process comprises obtaining a Hall effect voltage 3072, for example, through a Hall effect sensor located at the distal tip of the anvil 3013. The Hall effect voltage 3072 is provided to an analog to digital convertor 3074 and converted into a digital signal. The digital signal is provided to a processor, such as, for example, the primary processor 2006. The primary processor 2006 applies calibrates 3076 the curve input of the Hall effect voltage 3072 signal. A strain gauge 3078, such as, for example, a micro-strain gauge, is configured to measure one or more parameters of the end effector 3011, such as, for example, the amplitude of the strain exerted on the anvil 3013 during a clamping operation. The measured strain is converted 3080 to a digital signal and provided to the processor, such as, for example, the primary processor 2006. The load sensor 3082 measures the clamping force of the anvil 3013 against the staple cartridge 3021. The measured clamping force is converted 3084 to a digital signal and provided to the processor, such as for example, the primary processor 2006. The primary processor 2006 uses one or more algorithms and/or lookup tables to adjust the Hall effect voltage 3072 in response to the strain measured by the strain gauge 3078 and the clamping force measured by the load sensor 3082 to reflect the true thickness and fullness of the bite of tissue clamped by the anvil 3013 and the staple cartridge 3021. The adjusted thickness is displayed 3026 to an operator by, for example, a display 2026 embedded in the surgical instrument 10.

[0201] FIG. 60 illustrates one aspect of an end effector

3100 comprising a first sensor 3108a and a second sensor 3108b. The end effector 3100 is similar to the end effector 3011. The end effector 3100 comprises an anvil, or anvil, 3102 pivotally coupled to a jaw member 3104.

⁵ The jaw member 3104 is configured to receive a staple cartridge 3106 therein. The end effector 3100 comprises a first sensor 3108a coupled to the anvil 3102. The first sensor 3108a is configured to measure one or more parameters of the end effector 3100, such as, for example,

¹⁰ the gap 3110 between the anvil 3102 and the staple cartridge 3106. The gap 3110 may correspond to, for example, a thickness of tissue clamped between the anvil 3102 and the staple cartridge 3106. The first sensor 3108a may comprise any suitable sensor for measuring one or

¹⁵ more parameters of the end effector. For example, in various aspects, the first sensor 3108a may comprise a magnetic sensor, such as a Hall effect sensor, a strain gauge, a pressure sensor, an inductive sensor, such as an eddy current sensor, a resistive sensor, a capacitive ²⁰ sensor, an optical sensor, and/or any other suitable sensor.

[0202] In some aspects, the end effector 3100 comprises a second sensor 3108b. The second sensor 3108b is coupled to jaw member 3104 and/or the staple cartridge 3106. The second sensor 3108b is configured to

detect one or more parameters of the end effector 3100. For example, in some aspects, the second sensor 3108b is configured to detect one or more instrument conditions such as, for example, a color of the staple cartridge 3106 coupled to the jaw member 3104, a length of the staple cartridge 3106, a clamping condition of the end effector

3100, the number of uses/number of remaining uses of the end effector 3100 and/or the staple cartridge 3106, and/or any other suitable instrument condition. The second sensor 3108b may comprise any suitable sensor for

ond sensor 3108b may comprise any suitable sensor for detecting one or more instrument conditions, such as, for example, a magnetic sensor, such as a Hall effect sensor, a strain gauge, a pressure sensor, an inductive sensor, such as an eddy current sensor, a resistive sensor, a capacitive sensor, an optical sensor, and/or any

40 sor, a capacitive sensor, an optical sensor, and/or any other suitable sensor.
 (02031 In one aspect input from the second sensor.

[0203] In one aspect, input from the second sensor 3108b may be used to calibrate the input of the first sensor 3108a. The second sensor 3108b may be configured

⁴⁵ to detect one or more parameters of the staple cartridge 3106, such as, for example, the color and/or length of the staple cartridge 3106. The detected parameters, such as the color and/or the length of the staple cartridge 3106, may correspond to one or more properties of the car-

⁵⁰ tridge, such as, for example, the height of the cartridge deck, the thickness of tissue useable/optimal for the staple cartridge, and/or the pattern of the staples in the staple cartridge 3106. The known parameters of the staple cartridge 3106 may be used to adjust the thickness meas-⁵⁵ urement provided by the first sensor 3108a. For example, if the staple cartridge 3106 has a higher deck height, the thickness measurement provided by the first sensor 3108a may be reduced to compensate for the added deck height. The adjusted thickness may be displayed to an operator, for example, through a display 2026 coupled to the surgical instrument 10.

[0204] FIG. 61 illustrates one aspect of an end effector 3150 comprising a first sensor 3158 and a plurality of secondary sensors 3160a, 3160b. The end effector 3150 comprises an anvil, or anvil, 3152 and a jaw member 3154. The jaw member 3154 is configured to receive a staple cartridge 3156. The anvil 3152 is pivotally moveable with respect to the jaw member 3154 to clamp tissue between the anvil 3152 and the staple cartridge 3156. The anvil comprises a first sensor 3158. The first sensor 3158 is configured to detect one or more parameters of the end effector 3150, such as, for example, the gap 3110 between the anvil 3152 and the staple cartridge 3156. The gap 3110 may correspond to, for example, a thickness of tissue clamped between the anvil 3152 and the staple cartridge 3156. The first sensor 3158 may comprise any suitable sensor for measuring one or more parameters of the end effector. For example, in various aspects, the first sensor 3158 may comprise a magnetic sensor, such as a Hall effect sensor, a strain gauge, a pressure sensor, an inductive sensor, such as an eddy current sensor, a resistive sensor, a capacitive sensor, an optical sensor, and/or any other suitable sensor.

[0205] In some aspects, the end effector 3150 comprises a plurality of secondary sensors 3160a, 3160b. The secondary sensors 3160a, 3160b are configured to detect one or more parameters of the end effector 3150. For example, in some aspects, the secondary sensors 3160a, 3160b are configured to measure an amplitude of strain exerted on the anvil 3152 during a clamping procedure. In various aspects, the secondary sensors 3160a, 3160b may comprise a magnetic sensor, such as a Hall effect sensor, a strain gauge, a pressure sensor, an inductive sensor, such as an eddy current sensor, a resistive sensor, a capacitive sensor, an optical sensor, and/or any other suitable sensor. The secondary sensors 3160a, 3160b may be configured to measure one or more identical parameters at different locations of the anvil 3152, different parameters at identical locations on the anvil 3152, and/or different parameters at different locations on the anvil 3152.

[0206] FIG. 62 illustrates one aspect of an end effector 3200 comprising a plurality of sensors 3208a-3208d. The end effector 3200 comprises an anvil 3202 pivotally coupled to a jaw member 3204. The jaw member 3204 is configured to receive a staple cartridge 3206 therein. The anvil 3202 comprises a plurality of sensors 3208a-3208d thereon. The plurality of sensors 3208a-3208d is configured to detect one or more parameters of the end effector 3200, such as, for example, the anvil 3202. The plurality of sensors 3208a-3208d may comprise one or more identical sensors and/or different sensors. The plurality of sensors 3208a-3208d may comprise, for example, magnetic sensors, such as a Hall effect sensor, strain gauges, pressure sensors, inductive sensors, capacitive sensors, op-

tical sensors, and/or any other suitable sensors or combination thereof. For example, in one aspect, the plurality of sensors 3208a-3208d may comprise a plurality of strain gauges.

⁵ [0207] In one aspect, the plurality of sensors 3208a-3208d allows a robust tissue thickness sensing process to be implemented. By detecting various parameters along the length of the anvil 3202, the plurality of sensors 3208a-3208d allow a surgical instrument, such as, for

10 example, the surgical instrument 10, to calculate the tissue thickness in the jaws regardless of the bite, for example, a partial or full bite. In some aspects, the plurality of sensors 3208a-3208d comprises a plurality of strain gauges. The plurality of strain gauges is configured to

¹⁵ measure the strain at various points on the anvil 3202. The amplitude and/or the slope of the strain at each of the various points on the anvil 3202 can be used to determine the thickness of tissue in between the anvil 3202 and the staple cartridge 3206. The plurality of strain gaug-

²⁰ es may be configured to optimize maximum amplitude and/or slope differences based on clamping dynamics to determine thickness, tissue placement, and/or material properties of the tissue. Time based monitoring of the plurality of sensors 3208a-3208d during clamping allows

²⁵ a processor, such as, for example, the primary processor
 2006, to utilize algorithms and look-up tables to recognize
 tissue characteristics and clamping positions and dy namically adjust the end effector 3200 and/or tissue
 clamped between the anvil 3202 and the staple cartridge
 3206.

[0208] FIG. 63 is a logic diagram illustrating one aspect of a process 3220 for determining one or more tissue properties based on a plurality of sensors 3208a-3208d. In one aspect, a plurality of sensors 3208a-3208d generate 3222a-3222d a plurality of signals indicative of one or more parameters of the end effector 3200. The plurality of generated signals is converted 3224a-3224d to digital signals and provided to a processor. For example, in one aspect comprising a plurality of strain gauges, a plurality

40 of electronic μStrain (micro-strain) conversion circuits convert 3224a-3224d the strain gauge signals to digital signals. The digital signals are provided to a processor, such as, for example, the primary processor 2006. The primary processor 2006 determines 3226 one or more

⁴⁵ tissue characteristics based on the plurality of signals. The primary processor 2006 may determine the one or more tissue characteristics by applying an algorithm and/or a look-up table. The one or more tissue characteristics are displayed 3026 to an operator, for example,
⁵⁰ by a display 2026 embedded in the surgical instrument 10.

[0209] FIG. 64 illustrates one aspect of an end effector 3250 comprising a plurality of secondary sensors 3260a-3260d coupled to a jaw member 3254. The end effector 3250 comprises an anvil 3252 pivotally coupled to a jaw member 3254. The anvil 3252 is moveable relative to the jaw member 3254 to clamp one or more materials, such as, for example, a tissue section 3264, therebetween.

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The jaw member 3254 is configured to receive a staple cartridge 3256. A first sensor 3258 is coupled to the anvil 3252. The first sensor is configured to detect one or more parameters of the end effector 3150, such as, for example, the gap 3110 between the anvil 3252 and the staple cartridge 3256. The gap 3110 may correspond to, for example, a thickness of tissue clamped between the anvil 3252 and the staple cartridge 3256. The first sensor 3258 may comprise any suitable sensor for measuring one or more parameters of the end effector. For example, in various aspects, the first sensor 3258 may comprise a magnetic sensor, such as a Hall effect sensor, a strain gauge, a pressure sensor, an inductive sensor, such as an eddy current sensor, a resistive sensor, a capacitive sensor, an optical sensor, and/or any other suitable sensor.

[0210] A plurality of secondary sensors 3260a-3260d is coupled to the jaw member 3254. The plurality of secondary sensors 3260a-3260d may be formed integrally with the jaw member 3254 and/or the staple cartridge 3256. For example, in one aspect, the plurality of secondary sensors 3260a-3260d is disposed on an outer row of the staple cartridge 3256 (see FIG. 63). The plurality of secondary sensors 3260a-3260d are configured to detect one or more parameters of the end effector 3250 and/or a tissue section 3264 clamped between the anvil 3252 and the staple cartridge 3256. The plurality of secondary sensors 3260a-3260d may comprise any suitable sensors for detecting one or more parameters of the end effector 3250 and/or the tissue section 3264, such as, for example, magnetic sensors, such as a Hall effect sensor, strain gauges, pressure sensors, inductive sensors, such as an eddy current sensor, resistive sensors, capacitive sensors, optical sensors, and/or any other suitable sensors or combination thereof. The plurality of secondary sensors 3260a-3260d may comprise identical sensors and/or different sensors.

[0211] In some aspects, the plurality of secondary sensors 3260a-3260d comprises dual purpose sensors and tissue stabilizing elements. The plurality of secondary sensors 3260a-3260d comprise electrodes and/or sensing geometries configured to create a stabilized tissue condition when the plurality of secondary sensors 3260a-3260d are engaged with a tissue section 3264, such as, for example, during a clamping operation. In some aspects, one or more of the plurality of secondary sensors 3260a-3260d may be replaced with non-sensing tissue stabilizing elements. The secondary sensors 3260a-3260d create a stabilized tissue condition by controlling tissue flow, staple formation, and/or other treatment process.

[0212] FIG. 65 illustrates one aspect of a staple cartridge 3270 comprising a plurality of sensors 3272a-3272h formed integrally therein. The staple cartridge 3270 comprises a plurality of rows containing a plurality of holes for storing staples therein. One or more of the holes in the outer row 3278 are replaced with one of the plurality of sensors 3272a-3272h. A cutaway section 3274 is shown to illustrate a sensor 3272f coupled to a sensor wire 3276b. The sensor wires 3276a, 3276b may comprise a plurality of wires for coupling the plurality of sensors 3272a-3272h to one or more circuits of a surgical instrument, such as, for example, the surgical instrument 10. In some aspects, one or more of the plurality of sensors 3272a-3272h comprise dual purpose sensor and tissue stabilizing elements having electrodes and/or

- ¹⁰ sensing geometries configured to provide tissue stabilization. In some aspects, the plurality of sensors 3272a-3272h may be replaced with and/or co-populated with a plurality of tissue stabilizing elements. Tissue stabilization may be provided by, for example, controlling tissue
- ¹⁵ flow and/or staple formation during a clamping and/or stapling process. The plurality of sensors 3272a-3272h provide signals to one or more circuits of the surgical instrument 10 to enhance feedback of stapling performance and/or tissue thickness sensing.

20 [0213] FIG. 66 is a logic diagram illustrating one aspect of a process 3280 for determining one or more parameters of a tissue section 3264 clamped within an end effector, such as, for example, the end effector 3250 illustrated in FIG. 64. In one aspect, a first sensor 3258 is

- ²⁵ configured to detect one or more parameters of the end effector 3250 and/or a tissue section 3264 located between the anvil 3252 and the staple cartridge 3256. A first signal is generated 3282 by the first sensors 3258. The first signal is indicative of the one or more parameters
- ³⁰ detected by the first sensor 3258. One or more secondary sensors 3260 are configured to detect one or more parameters of the end effector 3250 and/or the tissue section 3264. The secondary sensors 3260 may be configured to detect the same parameters, additional parame-
- ³⁵ ters, or different parameters as the first sensor 3258. Secondary signals 3284 are generated by the secondary sensors 3260. The secondary signals 3284 are indicative of the one or more parameters detected by the secondary sensors 3260. The first signal and the secondary signals
- 40 are provided to a processor, such as, for example, a primary processor 2006. The primary processor 2006 adjusts 3286 the first signal generated by the first sensor 3258 based on input generated by the secondary sensors 3260. The adjusted signal may be indicative of, for ex-
- ⁴⁵ ample, the true thickness of a tissue section 3264 and the fullness of the bite. The adjusted signal is displayed 3026 to an operator by, for example, a display 2026 embedded in the surgical instrument 10.

[0214] FIG. 67 illustrates one aspect of an end effector
3350 comprising a magnetic sensor 3358 comprising a specific sampling rate to limit or eliminate false signals. The end effector 3350 comprises an anvil, or anvil, 3352 pivotably coupled to a jaw member 3354. The jaw member 3354 is configured to receive a staple cartridge 3356
therein. The staple cartridge 3356 contains a plurality of staples that may be delivered to a tissue section located between the anvil 3352 and the staple cartridge 3356. A magnetic sensor 3358 is coupled to the anvil 3352. The

magnetic sensor 3358 is configured to detect one or more parameters of the end effector 3350, such as, for example, the gap 3364 between the anvil 3352 and the staple cartridge 3356. The gap 3364 may correspond to the thickness of a material, such as, for example, a tissue section, and/or the fullness of a bite of material located between the anvil 3352 and the staple cartridge 3356. The magnetic sensor 3358 may comprise any suitable sensor for detecting one or more parameters of the end effector 3350, such as, for example, a magnetic sensor, such as a Hall effect sensor, a strain gauge, a pressure sensor, an inductive sensor, such as an eddy current sensor, a resistive sensor, a capacitive sensor, an optical sensor, and/or any other suitable sensor.

[0215] In one aspect, the magnetic sensor 3358 comprises a magnetic sensor configured to detect a magnetic field generated by an electromagnetic source 3360 coupled to the jaw member 3354 and/or the staple cartridge 3356. The electromagnetic source 3360 generates a magnetic field detected by the magnetic sensor 3358. The strength of the detected magnetic field may correspond to, for example, the thickness and/or fullness of a bite of tissue located between the anvil 3352 and the staple cartridge 3356. In some aspects, the electromagnetic source 3360 generates a signal at a known frequency, such as, for example, 1 MHz. In other aspects, the signal generated by the electromagnetic source 3360 may be adjustable based on, for example, the type of staple cartridge 3356 installed in the jaw member 3354, one or more additional sensor, an algorithm, and/or one or more parameters.

[0216] In one aspect, a signal processor 3362 is coupled to the end effector 3350, such as, for example, the anvil 3352. The signal processor 3362 is configured to process the signal generated by the magnetic sensor 3358 to eliminate false signals and to boost the input from the magnetic sensor 3358. In some aspects, the signal processor 3362 may be located separately from the end effector 3350, such as, for example, in the handle assembly 14 of a surgical instrument 10. In some aspects, the signal processor 3362 is formed integrally with and/or comprises an algorithm executed by a general processor, such as, for example, the primary processor 2006. The signal processor 3362 is configured to process the signal from the magnetic sensor 3358 at a frequency substantially equal to the frequency of the signal generated by the electromagnetic source 3360. For example, in one aspect, the electromagnetic source 3360 generates a signal at a frequency of 1MHz. The signal is detected by the magnetic sensor 3358. The magnetic sensor 3358 generates a signal indicative of the detected magnetic field which is provided to the signal processor 3362. The signal is processed by the signal processor 3362 at a frequency of 1MHz to eliminate false signals. The processed signal is provided to a processor, such as, for example, the primary processor 2006. The primary processor 2006 correlates the received signal to one or more parameters of the end effector 3350, such as, for example, the gap 3364 between the anvil 3352 and the staple cartridge 3356.

[0217] FIG. 68 is a logic diagram illustrating one aspect of a process 3370 for generating a thickness measure⁵ ment for a tissue section located between an anvil and a staple cartridge of an end effector, such as, for example, the end effector 3350 illustrated in FIG. 45. In one aspect of the process 3370, a signal is generated 3372 by a modulated electromagnetic source 3360. The generated

¹⁰ signal may comprise, for example, a 1MHz signal. A magnetic sensor 3358 is configured to detect 3374 the signal generated by the electromagnetic source 3360. The magnetic sensor 3358 generates a signal indicative of the detected magnetic field and provides the signal to a

¹⁵ signal processor 3362. The signal processor 3362 processes 3376 the signal to remove noise, false signals, and/or to boost the signal. The processed signal is provided to an analog-to-digital convertor for conversion 3378 to a digital signal. Calibration 3380 of the digital

 signal may be performed, for example, by application of a calibration curve input algorithm and/or look-up table. The processes 3376, conversion 3378, and calibration 3380 may be performed by one or more circuits. The calibrated signal is displayed 3026 to a user by, for ex ample, a display 2026 formed integrally with the surgical instrument 10.

[0218] FIGS. 69A and 69B illustrate one aspect of an end effector 3800 comprising a pressure sensor. The end effector 3800 comprises an anvil, or anvil, 3802 pivotally 30 coupled to a jaw member 3804. The jaw member 3804 is configured to receive a staple cartridge 3806 therein. The staple cartridge 3806 comprises a plurality of staples. A first sensor 3808 is coupled to the anvil 3802 at a distal tip. The first sensor 3808 is configured to detect 35 one or more parameters of the end effector, such as, for example, the distance, or gap 3814, between the anvil 3802 and the staple cartridge 3806. The first sensor 3808 may comprise any suitable sensor, such as, for example, a magnetic sensor. A magnet 3810 may be coupled to 40 the jaw member 3804 and/or the staple cartridge 3806

to provide a magnetic signal to the magnetic sensor. [0219] In some aspects, the end effector 3800 comprises a second sensor 3812. The second sensor 3812 is configured to detect one or more parameters of the

end effector 3800 and/or a tissue section located therebetween. The second sensor 3812 may comprise any suitable sensor, such as, for example, one or more pressure sensors. The second sensor 3812 may be coupled to the anvil 3802, the jaw member 3804, and/or the staple
cartridge 3806. A signal from the second sensor 3812

may be used to adjust the measurement of the first sensor
3808 to adjust the reading of the first sensor to accurately
represent proximal and/or distal positioned partial bites
true compressed tissue thickness. In some aspects, the
second sensor 3812 may be surrogate with respect to
the first sensor 3808.

[0220] In some aspects, the second sensor 3812 may comprise, for example, a single continuous pressure

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sensing film and/or an array of pressure sensing films. The second sensor 3812 is coupled to the deck of the staple cartridge 3806 along the central axis covering, for example, a slot 3816 configured to receive a cutting and/or staple deployment member. The second sensor 3812 provides signals indicate of the amplitude of pressure applied by the tissue during a clamping procedure. During firing of the cutting and/or deployment member, the signal from the second sensor 3812 may be severed, for example, by cutting electrical connections between the second sensor 3812 and one or more circuits. In some aspects, a severed circuit of the second sensor 3812 may be indicative of a spent staple cartridge 3806. In other aspects, the second sensor 3812 may be positioned such that deployment of a cutting and/or deployment member does not sever the connection to the second sensor 3812. [0221] FIG. 70 illustrates one aspect of an end effector 3850 comprising a second sensor 3862 located between a staple cartridge 3806 and a jaw member 3804. The end effector 3850 comprises an anvil, or anvil, 3852 pivotally coupled to a jaw member 3854. The jaw member 3854 is configured to receive a staple cartridge 3856 therein. A first sensor 3858 is coupled to the anvil 3852 at a distal tip. The first sensor 3858 is configured to detect one or more parameters of the end effector 3850, such as, for example, the distance, or gap 3864, between the anvil 3852 and the staple cartridge 3856. The first sensor 3858 may comprise any suitable sensor, such as, for example, a magnetic sensor. A magnet 3860 may be coupled to the jaw member 3854 and/or the staple cartridge 3856 to provide a magnetic signal to the magnetic sensor. In some aspects, the end effector 3850 comprises a second sensor 3862 similar in all respect to the second sensor 3812 of FIGS. 69A-69B, except that it is located between the staple cartridge 3856 and the jaw member 3854.

[0222] FIG. 71 is a logic diagram illustrating one aspect of a process 3870 for determining and displaying the thickness of a tissue section clamped in an end effector 3800 or 3850, according to FIGS. 69A-69B or FIG. 70. The process comprises obtaining a Hall effect voltage 3872, for example, through a Hall effect sensor located at the distal tip of the anvil 3802. The Hall effect voltage 3872 is proved to an analog to digital converter 3876 and converted into a digital signal. The digital signal is provided to a process, such as for example the primary processor 2006. The primary processor 2006 calibrates 3874 the curve input of the Hall effect voltage 3872 signal. Pressure sensors, such as for example, second sensor 3812, is configured to measure 3880 one or more parameters of, for example, the end effector 3800, such as for example the amount of pressure being exerted by the anvil 3802 on the tissue clamped in the end effector 3800. In some aspects the pressure sensors may comprise a single continuous pressure sensing film and/or array of pressure sensing films. The pressure sensors may thus be operable determine variations in the measure pressure at different locations between the proximal and distal ends of the end effector 3800. The measured pressure is provided to the processor, such as for example the primary processor 2006. The primary processor 2006 uses one or more algorithms and/or lookup tables to adjust 3882 the Hall effect voltage 3872 in response to the pressure measured 3880 by the pressure sensors to more accurately reflect the thickness of the tissue clamped between, for example, the anvil 3802 and the staple cartridge 3806. The adjusted thickness is displayed 3878 to an operator by, for example, a display 2026 embedded in the surgical instrument 10.

[0223] FIG. 72 illustrates one aspect of an end effector 3900 comprising a plurality of second sensors 3192a-3192b located between a staple cartridge 3906 and an elongated channel 3904. The end effector 3900 compris-

¹⁵ es an anvil 3902 pivotally coupled to a jaw member or elongated channel 3904. The elongated channel 3904 is configured to receive a staple cartridge 3906 therein. The anvil 3902 further comprises a first sensor 3908 located in the distal tip. The first sensor 3908 is configured to

20 detect one or more parameters of the end effector 3900, such as, for example, the distance, or gap, between the anvil 3902 and the staple cartridge 3906. The first sensor 3908 may comprise any suitable sensor, such as, for example, a magnetic sensor. A magnet 3910 may be 25 coupled to the elongated channel 3904 and/or the staple cartridge 3906 to provide a magnetic signal to the first sensor 3908. In some aspects, the end effector 3900 comprises a plurality of second sensors 3912a-3912c located between the staple cartridge 3906 and the elon-30 gated channel 3904. The second sensors 3912a-3912c may comprise any suitable sensors, such as for instance piezo-resistive pressure film strips. In some aspects, the second sensors 3912a-3912c may be uniformly distrib-

uted between the distal and proximal ends of the end ³⁵ effector 3900. [0224] In some aspects, signals from the second sen-

sors 3912a-3912c may be used to adjust the measurement of the first sensor 3908. For instance, the signals from the second sensors 3912a-3912c may be used to
adjust the reading of the first sensor 3908 to accurately represent the gap between the anvil 3902 and the staple cartridge 3906, which may vary between the distal and proximal ends of the end effector 3900, depending on the location and/or density of tissue 3920 between the

45 anvil 3902 and the staple cartridge 3906. FIG. 11 illustrates an example of a partial bite of tissue 3920. As illustrated for purposes of this example, the tissue is located only in the proximal area of the end effector 3900, creating a high pressure 3918 area near the proximal 50 area of the end effector 3900 and a corresponding low pressure 3916 area near the distal end of the end effector. [0225] FIGS. 73A and 73B further illustrate the effect of a full versus partial bite of tissue 3920. FIG. 73A illustrates the end effector 3900 with a full bite of tissue 3920, 55 where the tissue 3920 is of uniform density. With a full bite of tissue 3920 of uniform density, the measured first gap 3914a at the distal tip of the end effector 3900 may be approximately the same as the measured second gap

3922a in the middle or proximal end of the end effector 3900. For example, the first gap 3914a may measure 2.4 mm, and the second gap may measure 2.3 mm. FIG. 73B illustrates an end effector 3900 with a partial bite of tissue 3920, or alternatively a full bit of tissue 3920 of non-uniform density. In this case, the first gap 3914b will measure less than the second gap 3922b measured at the thickest or densest portion of the tissue 3920. For example, the first gap may measure 1.0 mm, while the second gap may measure 1.9 mm. In the conditions illustrated in FIGS. 73A-73B, signals from the second sensors 3912a-3912c, such as for instance measured pressure at different points along the length of the end effector 3900, may be employed by the instrument to determine tissue 3920 placement and/or material properties of the tissue 3920. The instrument may further be operable to use measured pressure over time to recognize tissue characteristics and tissue position, and dynamically adjust tissue thickness measurements.

[0226] FIG. 74 illustrates an aspect of an end effector 4050 that is configured to determine the location of a cutting member or knife 4062. The end effector 4050 comprises an anvil 4052 pivotally coupled to a j aw member or elongated channel 4054. The elongated channel 4054 is configured to receive a staple cartridge 4056 therein. The staple cartridge 4056 further comprises a slot (not shown) and a cutting member or knife 4062 located therein. The knife 4062 is operably coupled to a knife bar 4064. The knife bar 4064 is operable to move the knife 4062 from the proximal end of the slot to the distal end. The end effector 4050 may further comprise an optical sensor 4060 located near the proximal end of the slot. The optical sensor may be coupled to a processor, such as for instance the primary processor 2006. The optical sensor 4060 may be operable to emit an optical signal towards the knife bar 4064. The knife bar 4064 may further comprise a code strip 4066 along its length. The code strip 4066 may comprise cut-outs, notches, reflective pieces, or any other configuration that is optically readable. The code strip 4066 is placed such that the optical signal from the optical sensor 4060 will reflect off or through the code strip 4066. As the knife 4062 moves and knife bar 4064 moves 4068 along the slot 4058, the optical sensor 4060 will detect the reflection of the emitted optical signal coupled to the code strip 4066. The optical sensor 4060 may be operable to communicate the detected signal to the primary processor 2006. The primary processor 2006 may be configured to use the detected signal to determine the position of the knife 4062. The position of the knife 4062 may be sensed more precisely by designing the code strip 4066 such that the detected optical signal has a gradual rise and fall.

[0227] FIG. 75 illustrates an example of the code strip 4066 in operation with red LEDs 4070 and infrared LEDs 4072. For purposes of this example only, the code strip 4066 comprises cut-outs. As the code strip 4066 moves 4068, the light emitted by the red LEDs 4070 will be interrupted as the cut-outs passed before it. The infrared

LEDs 4072 will therefore detect the motion of the code strip 4066, and therefore, by extension, the motion of the knife 4062.

- **[0228]** FIG. 76 depicts a partial view of the end effector 300 of the surgical instrument 10. In the example form depicted in FIG. 76, the end effector 300 comprises a staple cartridge 1100 which is similar in many respects to the surgical staple cartridge 304 (FIG. 15). Several parts of the end effector 300 are omitted to enable a clear-
- ¹⁰ er understanding of the present disclosure. In certain instances, the end effector 300 may include a first jaw such as, for example, the anvil 306 (FIG. 20) and a second jaw such as, for example, the elongated channel 198 (FIG. 14). In certain instances, as described above, the

elongated channel 198 may accommodate a staple cartridge such as, for example, the surgical staple cartridge 304 or the staple cartridge 1100, for example. At least one of the elongated channel 198 and the anvil 306 may be movable relative to the other one of the elongated
channel 198 and the anvil 306 to capture tissue between the staple cartridge 1100 and the anvil 306. Various actuation assemblies are described herein to facilitation motion of the elongated channel 198 and/or the anvil 306 between an open configuration (FIG. 1) and a closed configuration (FIG. 77), for example.

configuration (FIG. 77), for example. [0229] In certain instances, as described above, the Ebeam 178 can be advanced distally to deploy the staples 191 into the captured tissue and/or advance the cutting edge 182 between a plurality of positions to engage and 30 cut the captured tissue. As illustrated in FIG. 76, the cutting edge 182 can be advanced distally along a path defined by the slot 193, for example. In certain instances, the cutting edge 182 can be advanced from a proximal portion 1103 of the staple cartridge 1100 to a distal por-35 tion 1105 of the staple cartridge 1100 to cut the captured tissue. In certain instances, the cutting edge 182 can be retracted proximally from the distal portion 1105 to the proximal portion 1103 by retraction of the E-beam 178 proximally, for example.

⁴⁰ **[0230]** In certain instances, the cutting edge 182 can be employed to cut tissue captured by the end effector 300 in multiple procedures. The reader will appreciate that repetitive use of the cutting edge 182 may affect the sharpness of the cutting edge 182. The reader will also

45 appreciate that as the sharpness of the cutting edge 182 decreases, the force required to cut the captured tissue with the cutting edge 182 may increase. Referring to FIGS. 78-83, in certain instances, the surgical instrument 10 may comprise a circuit 1106 (FIG. 78) for monitoring 50 the sharpness of the cutting edge 182 during, before, and/or after operation of the surgical instrument 10 in a surgical procedure, for example. In certain instances, the circuit 1106 can be employed to test the sharpness of the cutting edge 182 prior to utilizing the cutting edge 182 55 to cut the captured tissue. In certain instances, the circuit 1106 can be employed to test the sharpness of the cutting edge 182 after the cutting edge 182 has been used to cut the captured tissue. In certain instances, the circuit

1106 can be employed to test the sharpness of the cutting edge 182 prior to and after the cutting edge 182 is used to cut the captured tissue. In certain instances, the circuit 1106 can be employed to test the sharpness of the cutting edge 182 at the proximal portion 1103 and/or at the distal portion 1105.

[0231] Referring to FIGS. 78-83, the circuit 1106 may include one or more sensors such as, for example, an optical sensor 1108; the optical sensor 1108 of the circuit 1106 can be employed to test the reflective ability of the cutting edge 182, for example. In certain instances, the ability of the cutting edge 182 to reflect light may correlate with the sharpness of the cutting edge 182. In other words, a decrease in the sharpness of the cutting edge 182 may result in a decrease in the ability of the cutting edge 182 to reflect the light. Accordingly, in certain instances, the dullness of the cutting edge 182 can be evaluated by monitoring the intensity of the light reflected from the cutting edge 182, for example. In certain instances, the optical sensor 1108 may define a light sensing region. The optical sensor 1108 can be oriented such that the optical sensing region is disposed in the path of the cutting edge 182, for example. The optical sensor 1108 may be employed to sense the light reflected from the cutting edge 182 while the cutting edge 182 is in the optical sensing region, for example. A decrease in intensity of the reflected light beyond a threshold can indicate that the sharpness of the cutting edge 182 has decreased beyond an acceptable level.

[0232] Referring again to FIGS. 78-83, the circuit 1106 may include one or more lights sources such as, for example, a light source 1110. In certain instances, the circuit 1106 may include a controller 1112 ("microcontroller") which may be operably coupled to the optical sensor 1108, as illustrated in FIGS. 78-83. In certain instances, the controller 1112 may include a processor 1114 ("microprocessor") and one or more computer readable mediums or memory 1116 ("memory units"). In certain instances, the memory 1116 may store various program instructions, which when executed may cause the processor 1114 to perform a plurality of functions and/or calculations described herein. In certain instances, the memory 1116 may be coupled to the processor 1114, for example. A power source 1118 can be configured to supply power to the controller 1112, the optical sensors 1108, and/or the light sources 1110, for example. In certain instances, the power source 1118 may comprise a battery (or "battery pack" or "power pack"), such as a Li ion battery, for example. In certain instances, the battery pack may be configured to be releasably mounted to the handle assembly 14 for supplying power to the surgical instrument 10. A number of battery cells connected in series may be used as the power source 4428. In certain instances, the power source 1118 may be replaceable and/or rechargeable, for example.

[0233] The controller 1112 and/or other controllers of the present disclosure may be implemented using integrated and/or discrete hardware elements, software elements, and/or a combination of both. Examples of integrated hardware elements may include processors, microprocessors, controllers, integrated circuits, ASICs, PLDs, DSPs, FPGAs, logic gates, registers, semiconductor devices, chips, microchips, chip sets, controllers, SoC, and/or SIP. Examples of discrete hardware elements may include circuits and/or circuit elements such as logic gates, field effect transistors, bipolar transistors, resistors, capacitors, inductors, and/or relays. In certain

¹⁰ instances, the controller 1112 may include a hybrid circuit comprising discrete and integrated circuit elements or components on one or more substrates, for example. In certain instances, the controller 1112 and/or other controllers of the present disclosure may be a single core or

¹⁵ multicore controller LM4F230H5QR as described in connection with FIGS. 14-17B.
 [0234] In certain instances, the light source 1110 can

be employed to emit light which can be directed at the cutting edge 182 in the optical sensing region, for exam-

- 20 ple. The optical sensor 1108 may be employed to measure the intensity of the light reflected from the cutting edge 182 while in the optical sensing region in response to exposure to the light emitted by the light source 1110. In certain instances, the processor 1114 may receive one
- or more values of the measured intensity of the reflected light and may store the one or more values of the measured intensity of the reflected light on the memory 1116, for example. The stored values can be detected and/or recorded before, after, and/or during a plurality of surgical
 procedures performed by the surgical instrument 10, for

example. **[0235]** In certain instances, the processor 1114 may compare the measured intensity of the reflected light to a predefined threshold values that may be stored on the 35 memory 1116, for example. In certain instances, the controller 1112 may conclude that the sharpness of the cutting edge 182 has dropped below an acceptable level if the measured light intensity exceeds the predefined threshold value by 1%, 5%, 10%, 25%, 50%, 100% 40 and/or more than 100%, for example. In certain instances, the processor 1114 can be employed to detect a decreasing trend in the stored values of the measured intensity of the light reflected from the cutting edge 182 while in the optical sensing region.

45 [0236] In certain instances, the surgical instrument 10 may include one or more feedback systems such as, for example, the feedback system 1120. In certain instances, the processor 1114 can employ the feedback system 1120 to alert a user if the measured light intensity of the 50 light reflected from cutting edge 182 while in the optical sensing region is beyond the stored threshold value, for example. In certain instances, the feedback system 1120 may comprise one or more visual feedback systems such as display screens, backlights, and/or LEDs, for example. 55 In certain instances, the feedback system 1120 may comprise one or more audio feedback systems such as speakers and/or buzzers, for example. In certain instances, the feedback system 1120 may comprise one or more

haptic feedback systems, for example. In certain instances, the feedback system 1120 may comprise combinations of visual, audio, and/or tactile feedback systems, for example.

[0237] In certain instances, the surgical instrument 10 may comprise a firing lockout mechanism 1122 which can be employed to prevent advancement of the cutting edge 182. Various suitable firing lockout mechanisms are described in greater detail in U.S. Patent Application Publication No. 2014/0001231, entitled FIRING SYS-TEM LOCKOUT ARRANGEMENTS FOR SURGICAL INSTRUMENTS, which is herein incorporated by reference in its entirety. In certain instances, as illustrated in FIG. 78, the processor 1114 can be operably coupled to the firing lockout mechanism 1122; the processor 1114 may employ the firing lockout mechanism 1122 to prevent advancement of the cutting edge 182 in the event it is determined that the measured intensity of the light reflected from the cutting edge 182 is beyond the stored threshold, for example. In other words, the processor 1114 may activate the firing lockout mechanism 1122 if the cutting edge is not sufficiently sharp to cut the tissue captured by the end effector 300.

[0238] In certain instances, the optical sensor 1108 and the light source 1110 can be housed at a distal portion of the interchangeable shaft assembly 200. In certain instances, the sharpness of cutting edge 182 can be evaluated by the optical sensor 1108, as described above, prior to transitioning the cutting edge 182 into the end effector 300. The firing bar 172 (FIG. 14) may advance the cutting edge 182 through the optical sensing region defined by the optical sensor 1108 while the cutting edge 182 is in the interchangeable shaft assembly 200 and prior to entering the end effector 300, for example. In certain instances, the sharpness of cutting edge 182 can be evaluated by the optical sensor 1108 after retracting the cutting edge 182 proximally from the end effector 300. The firing bar 172 (FIG. 14) may retract the cutting edge 182 through the optical sensing region defined by the optical sensor 1108 after retracting the cutting edge 182 from the end effector 300 into the interchangeable shaft assembly 200, for example.

[0239] In certain instances, the optical sensor 1108 and the light source 1110 can be housed at a proximal portion of the end effector 300 which can be proximal to the staple cartridge 1100, for example. The sharpness of cutting edge 182 can be evaluated by the optical sensor 1108 after transitioning the cutting edge 182 into the end effector 300 but prior to engaging the staple cartridge 1100, for example. In certain instances, the firing bar 172 (FIG. 14) may advance the cutting edge 182 through the optical sensing region defined by the optical sensor 1108 while the cutting edge 182 is in the end effector 300 but prior to engaging the staple cartridge 1100, for example. [0240] In various instances, the sharpness of cutting edge 182 can be evaluated by the optical sensor 1108 as the cutting edge 182 is advanced by the firing bar 172 through the slot 193. As illustrated in FIGS. 78-83, the

optical sensor 1108 and the light source 1110 can be housed at the proximal portion 1103 of the staple cartridge 1100, for example; and the sharpness of cutting edge 182 can be evaluated by the optical sensor 1108 at the proximal portion 1103, for example. The firing bar

172 (FIG. 14) may advance the cutting edge 182 through the optical sensing region defined by the optical sensor 1108 at the proximal portion 1103 before the cutting edge 182 engages tissue captured between the staple car-

10 tridge 1100 and the anvil 306, for example. In certain instances, as illustrated in FIGS. 78-83, the optical sensor 1108 and the light source 1110 can be housed at the distal portion 1105 of the staple cartridge 1100, for example. The sharpness of cutting edge 182 can be eval-

15 uated by the optical sensor 1108 at the distal portion 1105. In certain instances, the firing bar 172 (FIG. 14) may advance the cutting edge 182 through the optical sensing region defined by the optical sensor 1108 at the distal portion 1105 after the cutting edge 182 has passed 20 through the tissue captured between the staple cartridge

1100 and the anvil 306, for example. [0241] Referring again to FIG. 76, the staple cartridge 1100 may comprise a plurality of optical sensors 1108 and a plurality of corresponding light sources 1110, for 25 example. In certain instances, a pair of the optical sensor 1108 and the light source 1110 can be housed at the proximal portion 1103 of the staple cartridge 1100, for example; and a pair of the optical sensor 1108 and the light source 1110 can be housed at the distal portion 1105 30 of the staple cartridge 1100, for example. In such instances, the sharpness of the cutting edge 182 can be evaluated a first time at the proximal portion 1103 prior to engaging the tissue, for example, and a second time at the distal portion 1105 after passing through the captured

tissue, for example. [0242] The reader will appreciate that an optical sensor 1108 may evaluate the sharpness of the cutting edge 182 a plurality of times during a surgical procedure. For example, the sharpness of the cutting edge can be eval-40 uated a first time during advancement of the cutting edge 182 through the slot 193 in a firing stroke, and a second time during retraction of the cutting edge 182 through the slot 193 in a return stroke, for example. In other words,

the light reflected from the cutting edge 182 can be meas-45 ured by the optical sensor 1108 once as the cutting edge is advanced through the optical sensing region, and once as the cutting edge 182 is retracted through the optical sensing region, for example.

[0243] The reader will appreciate that the processor 50 1114 may receive a plurality of readings of the intensity of the light reflected from the cutting edge 182 from one or more of the optical sensors 1108. In certain instances, the processor 1114 may be configured to discard outliers and calculate an average reading from the plurality of 55 readings, for example. In certain instances, the average reading can be compared to a threshold stored in the memory 1116, for example. In certain instances, the processor 1114 may be configured to alert a user through

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the feedback system 1120 and/or activate the firing lockout mechanism 1122 if it is determined that the calculated average reading is beyond the threshold stored in the memory 1116, for example.

[0244] In certain instances, as illustrated in FIGS. 77, 79, and 80, a pair of the optical sensor 1108 and the light source 1110 can be positioned on opposite sides of the staple cartridge 1100. In other words, the optical sensor 1108 can be positioned on a first side 1124 of the slot 193, for example, and the light source 1110 can be positioned on a second side 1126, opposite the first side 1124, of the slot 193, for example. In certain instances, the pair of the optical sensor 1108 and the light source 1110 can be substantially disposed in a plane transecting the staple cartridge 1100, as illustrated in FIG. 77. The pair of the optical sensor 1108 and the light source 1110 can be oriented to define an optical sensing region that is positioned, or at least substantially positioned, on the plane transecting the staple cartridge 1100, for example. Alternatively, the pair of the optical sensor 1108 and the light source 1110 can be oriented to define an optical sensing region that is positioned proximal to the plane transecting the staple cartridge 1100, for example, as illustrated in FIG. 80.

[0245] In certain instances, a pair of the optical sensor 1108 and the light source 1110 can be positioned on a same side of the staple cartridge 1100. In other words, as illustrated in FIG. 81, the pair of the optical sensor 1108 and the light source 1110 can be positioned on a first side of the cutting edge 182, e.g. the side 1128, as the cutting edge 182 is advanced through the slot 193. In such instances, the light source 1110 can be oriented to direct light at the side 1128 of the cutting edge 182; and the intensity of the light reflected from the side 1128, as measured by the optical sensor 1108, may represent the sharpness of the side 1128.

[0246] In certain instances, as illustrated in FIG. 82, a second pair of the optical sensor 1108 and the light source 1110 can be positioned on a second side of the cutting edge 182, e.g. the side 1130, for example. The second pair can be employed to evaluate the sharpness of the side 1130. For example, the light source 1110 of the second pair can be oriented to direct light at the side 1130 of the cutting edge 182; and the intensity of the light reflected from the side 1130, as measured by the optical sensor 1108 of the second pair, may represent the sharpness of the side 1130. In certain instances, the processor can be configured to assess the sharpness of the cutting edge 182 based upon the measured intensities of the light reflected from the sides 1128 and 1130 of the cutting edge 182, for example.

[0247] In certain instances, as illustrated in FIG. 77, a pair of the optical sensor 1108 and the light source 1110 can be housed at the distal portion 1105 of the staple cartridge 1100. As illustrated in FIG. 81, the optical sensor 1108 can be positioned, or at least substantially positioned, on an axis LL which extends longitudinally along the path of the cutting edge 182 through the slot 193, for

example. In addition, the light source 1110 can be positioned distal to the cutting edge 182 and oriented to direct light at the cutting edge 182 as the cutting edge is advanced toward the light source 1110, for example. Furthermore, the optical sensor 1108 can be positioned, or at least substantially positioned, along an axis AA that intersects the axis LL, as illustrated in FIG. 81. In certain

instances, the axis AA may be perpendicular to the axis LL, for example. In any event, the optical sensor 1108 can be oriented to define an optical sensing region at the intersection of the axis LL and the axis AA, for example.

[0248] The reader will appreciate that the position, orientation and/or number of optical sensors and corresponding light sources described herein in connection

¹⁵ with the surgical instrument 10 are example aspects intended for illustration purposes. Various other arrangements of optical sensors and light sources can be employed by the present disclosure to evaluate the sharpness of the cutting edge 182.

20 [0249] The reader will appreciate that advancement of the cutting edge 182 through the tissue captured by the end effector 300 may cause the cutting edge to collect tissue debris and/or bodily fluids during each firing of the surgical instrument 10. Such debris may interfere with

the ability of the circuit 1106 to accurately evaluate the sharpness of the cutting edge 182. In certain instances, the surgical instrument 10 can be equipped with one or more cleaning mechanisms which can be employed to clean the cutting edge 182 prior to evaluating the sharpness of the cutting edge 182, for example.

[0250] Referring to FIG. 76, in certain instances, the staple cartridge 1100 may include a first pair of the optical sensor 1108 and the light source 1110, which can be housed in the proximal portion 1103 of the staple cartridge 1100, for example. Furthermore, as illustrated in FIG. 76, the staple cartridge 1100 may include a first pair of the cleaning members 1132, which can be housed in the proximal portion 1103 on opposite sides of the slot

193. The first pair of the cleaning members 1132 can be
positioned distal to the first pair of the optical sensor 1108
and the light source 1110, for example. As illustrated in
FIG. 76, the staple cartridge 1100 may include a second
pair of the optical sensor 1108 and the light source 1110,
which can be housed in the distal portion 1105 of the

staple cartridge 1100, for example. As illustrated in FIG.
76, the staple cartridge 1100 may include a second pair of the cleaning members 1132, which can be housed in the distal portion 1105 on opposite sides of the slot 193. The second pair of the cleaning members 1132 can be
positioned proximal to the second pair of the optical sensor 1108 and the light source 1110.

[0251] Further to the above, as illustrated in FIG. 76, the cutting edge 182 may be advanced distally in a firing stroke to cut tissue captured by the end effector 300. As the cutting edge is advanced, a first evaluation of the sharpness of the cutting edge 182 can be performed by the first pair of the optical sensor 1108 and the light source 1110 prior to tissue engagement by the cutting edge 182,

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for example. A second evaluation of the sharpness of the cutting edge 182 can be performed by the second pair of the optical sensor 1108 and the light source 1110 after the cutting edge 182 has transected the captured tissue, for example. The cutting edge 182 may be advanced through the second pair of the cleaning members 1132 prior to the second evaluation of the sharpness of the cutting edge 182 to remove any debris collected by the cutting edge 182 during the transection of the captured tissue.

[0252] Further to the above, as illustrated in FIG. 76, the cutting edge 182 may be retracted proximally in a return stroke. As the cutting edge is retracted, a third evaluation of the sharpness of the cutting edge 182 can be performed by the first pair of the optical sensor 1108 and the light source 1110 during the return stroke. The cutting edge 182 may be retracted through the first pair of the cleaning members 1132 prior to the third evaluation of the sharpness of the cutting edge 182 to remove any debris collected by the cutting edge 182 during the transection of the captured tissue, for example.

[0253] In certain instances, one or more of the lights sources 1110 may comprise one or more optical fiber cables. In certain instances, one or more flex circuits 1134 can be employed to transmit energy from the power source 1118 to the optical sensors 1108 and/or the light sources 1110. In certain instances, the flex circuits 1134 may be configured to transmit one or more of the readings of the optical sensors 1108 to the controller 1112, for example.

[0254] Referring now to FIG. 84, a staple cartridge 4300 is depicted; the staple cartridge 4300 is similar in many respects to the surgical staple cartridge 304 (FIG. 14). For example, the staple cartridge 4300 can be employed with the end effector 300. In certain instances, as illustrated in FIG. 84, the staple cartridge 4300 may comprise a sharpness testing member 4302 which can be employed to test the sharpness of the cutting edge 182. In certain instances, the sharpness testing member 4302 can be attached to and/or integrated with the cartridge body 194 of the staple cartridge 4300, for example. In certain instances, the sharpness testing member 4302 can be disposed in the proximal portion 1103 of the staple cartridge 4300, for example. In certain instances, as illustrated in FIG. 84, the sharpness testing member 4302 can be disposed onto a cartridge deck 4304 of the staple cartridge 4300, for example.

[0255] In certain instances, as illustrated in FIG. 84, the sharpness testing member 4302 can extend across the slot 193 of the staple cartridge 4300 to bridge, or at least partially bridge, the gap defined by the slot 193, for example. In certain instances, the sharpness testing member 4302 may interrupt, or at least partially interrupt, the path of the cutting edge 182. The cutting edge 182 may engage, cut, and/or pass through the sharpness testing member 4302 as the cutting edge 182 is advanced during a firing stroke, for example. In certain instances, the cutting edge 182 may be configured to engage, cut,

and/or pass through the sharpness testing member 4302 prior to engaging tissue captured by the end effector 300 in a firing stroke, for example. In certain instances, the cutting edge 182 may be configured to engage the sharpness testing member 4302 at a proximal end 4306 of the sharpness testing member 4302, and exit and/or disengage the sharpness testing member 4302 at a distal end 4308 of the sharpness testing member 4302, for example. In certain instances, the cutting edge 182 can travel

¹⁰ and/or cut through the sharpness testing member 4302 a distance (D) between the proximal end 4306 and the distal end 4308, for example, as the cutting edge 182 is advanced during a firing stroke.

[0256] Referring primarily to FIGS. 84 and 85, the surgical instrument 10 may comprise a circuit 4310 for testing the sharpness of the cutting edge 182, for example.
In certain instances, the circuit 4310 can evaluate the sharpness of the cutting edge 182 by testing the ability of the cutting edge 182 to be advanced through the sharpness testing member 4302. For example, the circuit 4310 can be configured to observe the time period the cutting edge 182 takes to fully transect and/or completely pass through at least a predetermined portion of the sharpness testing member 4302. If the observed time period exceeds a predetermined threshold, the circuit 4310 may

conclude that the sharpness of the cutting edge 182 has dropped below an acceptable level, for example.

[0257] In certain instances, the circuit 4310 may include a controller 4312 ("microcontroller") which may include a processor 4314 ("microprocessor") and one or more computer readable mediums or memory 4316 units ("memory"). In certain instances, the memory 4316 may store various program instructions, which when executed may cause the processor 4314 to perform a plurality of functions and/or calculations described herein. In certain

instances, the memory 4316 may be coupled to the processor 4314, for example. A power source 4318 can be configured to supply power to the controller 4312, for example. In certain instances, the power source 4138

40 may comprise a battery (or "battery pack" or "power pack"), such as a Li ion battery, for example. In certain instances, the battery pack may be configured to be releasably mounted to the handle assembly 14. A number of battery cells connected in series may be used as the

⁴⁵ power source 4318. In certain instances, the power source 4318 may be replaceable and/or rechargeable, for example.

[0258] In certain instances, the controller 4313 can be operably coupled to the feedback system 1120 and/or the firing lockout mechanism 1122, for example.

[0259] Referring to FIGS. 84 and 85, the circuit 4310 may comprise one or more position sensors. Example position sensors and positioning systems suitable for use with the present disclosure are described in U.S. Patent
 ⁵⁵ Application Publication No. 2014/0263538, entitled SEN-SOR ARRANGEMENTS FOR ABSOLUTE POSITION-ING SYSTEM FOR SURGICAL INSTRUMENTS, which is herein incorporated by reference in its entirety. In cer-

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tain instances, the circuit 4310 may include a first position sensor 4320 and a second position sensor 4322. In certain instances, the first position sensor 4320 can be employed to detect a first position of the cutting edge 182 at the proximal end 4306 of the sharpness testing member 4302, for example; and the second position sensor 4322 can be employed to detect a second position of the cutting edge 182 at the distal end 4308 of the sharpness testing member 4302, for example.

[0260] In certain instances, the first and second position sensors 4320, 4322 can be employed to provide first and second position signals, respectively, to the controller 4312. It will be appreciated that the position signals may be analog signals or digital values based on the interface between the controller 4312 and the first and second position sensors 4320, 4322. In one aspect, the interface between the controller 4312 and the first and second position sensors 4320, 4322 can be a standard serial peripheral interface (SPI), and the position signals can be digital values representing the first and second positions of the cutting edge 182, as described above.

[0261] Further to the above, the processor 4314 may determine the time period between receiving the first position signal and receiving the second position signal. The determined time period may correspond to the time it takes the cutting edge 182 to advance through the sharpness testing member 4302 from the first position at the proximal end 4306 of the sharpness testing member 4302, for example, to the second position at the distal end 4308 of the sharpness testing member 4302, for example. In at least one example, the controller 4312 may include a time element which can be activated by the processor 4314 upon receipt of the first position signal, and deactivated upon receipt of the second position signal. The time period between the activation and deactivation of the time element may correspond to the time it takes the cutting edge 182 to advance from the first position to the second position, for example. The time element may comprise a real time clock, a processor configured to implement a time function, or any other suitable timing circuit.

[0262] In various instances, the controller 4312 can compare the time period it takes the cutting edge 182 to advance from the first position to the second position to a predefined threshold value to assess whether the sharpness of the cutting edge 182 has dropped below an acceptable level, for example. In certain instances, the controller 4312 may conclude that the sharpness of the cutting edge 182 has dropped below an acceptable level if the measured time period exceeds the predefined threshold value by 1%, 5%, 10%, 25%, 50%, 100% and/or more than 100%, for example.

[0263] Referring to FIG. 86, in various instances, an electric motor 4330 can drive the firing bar 172 (FIG. 14) to advance the cutting edge 182 during a firing stroke and/or to retract the cutting edge 182 during a return stroke, for example. A motor driver 4332 can control the electric motor 4330; and a controller such as, for exam-

ple, the controller 4312 can be in signal communication with the motor driver 4332. As the electric motor 4330 advances the cutting edge 182, the controller 4312 can determine the current drawn by the electric motor 4330,

⁵ for example. In such instances, the force required to advance the cutting edge 182 can correspond to the current drawn by the electric motor 4330, for example. Referring still to FIG. 86, the controller 4312 of the surgical instrument 10 can determine if the current drawn by the electric

¹⁰ motor 4330 increases during advancement of the cutting edge 182 and, if so, can calculate the percentage increase of the current.

[0264] In certain instances, the current drawn by the electric motor 4330 may increase significantly while the

¹⁵ cutting edge 182 is in contact with the sharpness testing member 4302 due to the resistance of the sharpness testing member 4302 to the cutting edge 182. For example, the current drawn by the electric motor 4330 may increase significantly as the cutting edge 182 engages,

²⁰ passes and/or cuts through the sharpness testing member 4302. The reader will appreciate that the resistance of the sharpness testing member 4302 to the cutting edge 182 depends, in part, on the sharpness of the cutting edge 182; and as the sharpness of the cutting edge 182

decreases from repetitive use, the resistance of the sharpness testing member 4302 to the cutting edge 182 will increase. Accordingly, the value of the percentage increase of the current drawn by the electric motor 4330 while the cutting edge is in contact with the sharpness
testing member 4302 can increase as the sharpness of the cutting edge 182 decreases from repetitive use, for example.

[0265] In certain instances, the determined value of the percentage increase of the current drawn by the electric motor 4330 can be the maximum detected percent-

tric motor 4330 can be the maximum detected percentage increase of the current drawn by the electric motor 4330. In various instances, the controller 4312 can compare the determined value of the percentage increase of the current drawn by the electric motor 4330 to a prede-

fined threshold value of the percentage increase of the current drawn by the electric motor 4330. If the determined value exceeds the predefined threshold value, the controller 4312 may conclude that the sharpness of the cutting edge 182 has dropped below an acceptable level,
 for example.

[0266] In certain instances, as illustrated in FIG. 86, the processor 4314 can be in communication with the feedback system 1120 and/or the firing lockout mechanism 1122, for example. In certain instances, the processor 4314 can employ the feedback system 1120 to alert a user if the determined value of the percentage increase of the current drawn by the electric motor 4330 exceeds the predefined threshold value, for example. In certain instances, the processor 4314 may employ the firing lockout mechanism 1122 to prevent advancement of the cutting edge 182 if the determined value of the percentage increase of the current drawn by the electric motor 4330 exceeds the predefined threshold value, for the percentage increase of the current drawn by the electric motor 4330 exceeds the predefined threshold value of the percentage increase of the current drawn by the electric motor 4330 exceeds the predefined threshold value, for

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example.

[0267] In various instances, the controller 4312 can utilize an algorithm to determine the change in current drawn by the electric motor 4330. For example, a current sensor can detect the current drawn by the electric motor 4330 during the firing stroke. The current sensor can continually detect the current drawn by the electric motor and/or can intermittently detect the current draw by the electric motor. In various instances, the algorithm can compare the most recent current reading to the immediately proceeding current reading, for example. Additionally or alternatively, the algorithm can compare a sample reading within a time period X to a previous current reading. For example, the algorithm can compare the sample reading to a previous sample reading within a previous time period X, such as the immediately proceeding time period X, for example. In other instances, the algorithm can calculate the trending average of current drawn by the motor. The algorithm can calculate the average current draw during a time period X that includes the most recent current reading, for example, and can compare that average current draw to the average current draw during an immediately proceeding time period time X, for example.

[0268] Referring to FIG. 87, a method 4500 is depicted for evaluating the sharpness of the cutting edge 182 of the surgical instrument 10; and various responses are outlined in the event the sharpness of the cutting edge 182 drops to and/or below an alert threshold and/or a high severity threshold, for example. In various instances, a controller such as, for example, the controller 4312 can be configured to implement the method depicted in FIG. 85. In certain instances, the surgical instrument 10 may include a load cell 4334 (FIG. 86); as illustrated in FIG. 84, the controller 4312 may be in communication with the load cell 4334. In certain instances, the load cell 4334 may include a force sensor such as, for example, a strain gauge, which can be operably coupled to the firing bar 172, for example. In certain instances, the controller 4312 may employ the load cell 4334 to monitor the force (Fx) applied to the cutting edge 182 as the cutting edge 182 is advanced during a firing stroke.

[0269] Accordingly, when the knife firing is initiated 4502 the system checks 4504 the dullness of the cutting edge 182 of the knife by sensing a force Fx. The sensed force Fx is compared to a threshold force F1 and determines 4506 whether the sensed force Fx is greater than the threshold force F1. When the sensed force Fx is less than or equal to the threshold force F1, the process proceeds along NO branch and displays 4508 nothing and continues 4510 the knife firing process. When the sensed force Fx is greater than the threshold force F1, the process proceeds along YES branch and determines 4512 whether the sensed force Fx exceeds a high severity threshold force F2. When the sensed force Fx is less than or equal to the threshold F2, the process proceeds along NO branch and notifies 4514 the processor that the cutting edge 182 of the knife is dulling and the continues 4510 the knife firing process. When the sensed force Fx is greater than the threshold F2, the process proceeds along YES branch and notifies 4516 the processor that the cutting edge 182 of the knife is dulled and

⁵ the knife firing lockout is engaged. Subsequently, optionally, the processor may override 4518 the knife firing lockout and continues 4510 the knife firing process if the lockout is overridden.

[0270] Referring to FIG. 88, a method 4600 is depicted for determining whether a cutting edge such as, for example, the cutting edge 182 is sufficiently sharp to be employed in transecting a tissue of a particular tissue thickness that is captured by the end effector 300, for example. As described above, repetitive use of the cut-

¹⁵ ting edge 182 may dull or reduce the sharpness of the cutting edge 182 which may increase the force required for the cutting edge 182 to transect the captured tissue. In other words, the sharpness level of the cutting edge 182 can be defined by the force required for the cutting

20 edge 182 to transect the captured tissue, for example. The reader will appreciate that the force required for the cutting edge 182 to transect a captured tissue may also depend on the thickness of the captured tissue. In certain instances, the greater the thickness of the captured tis-

sue, the greater the force required for the cutting edge182 to transect the captured tissue at the same sharp-ness level, for example.

[0271] Accordingly, initially, the stapler clamps 4602 the tissue between the anvil and the jaw member. The system senses 4604 the tissue thickness Tx and initiates 4606 the knife firing process. Upon initiating the knife firing process, the system senses 4608 the load resistance from the clamped tissue and compares the sensed force Fx and senses thickness Tx against various thresholds and determines 4610 several outcomes based on

the evaluation. In one aspect, when the process determines 4610 whether the sensed tissue thickness Tx is within a first tissue thickness range defined between a first tissue thickness threshold T1 and a second tissue thickness threshold T2 AND the sensed force Fx is great-

thickness threshold T2 AND the sensed force Fx is greater than a first force threshold F1 AND the process determines 4610 whether the sensed tissue thickness Tx is within a second tissue thickness range defined between the second tissue thickness threshold T2 and a third tis-

45 sue thickness threshold T3 AND the sensed force Fx is greater than a second force threshold F2, the process proceeds along the YES branch and notifies 4612 or alerts the processor that the knife is dulling and then continues 4614 the knife firing process. Otherwise, the proc-50 ess proceeds along the NO branch and the does not notify 4616 the processor and continues the knife firing process. Generally, the process determines whether the sensed tissue thickness Tx is within a tissue thickness range defined between tissue thickness thresholds Tn 55 and Tn+1 AND the sensed force Fx is greater than a force threshold Tn, where n indicates a tissue thickness range. When the process determines 4610 that the sensed tissue thickness Tx is within a first tissue thick-

ness range defined between a first tissue thickness threshold T1 and a second tissue thickness threshold T2 AND the sensed force Fx is greater than a first force threshold F1 AND when the process determines 4610 that the sensed tissue thickness Tx is within a second tissue thickness range defined between the second tissue thickness threshold T2 and a third tissue thickness threshold T3 AND the sensed force Fx is greater than a second force threshold F2, the process continues.

[0272] In certain instances, the cutting edge 182 may be sufficiently sharp for transecting a captured tissue comprising a first thickness but may not be sufficiently sharp for transecting a captured tissue comprising a second thickness greater than the first thickness, for example. In certain instances, a sharpness level of the cutting edge 182, as defined by the force required for the cutting edge 182 to transect a captured tissue, may be adequate for transecting the captured tissue if the captured tissue comprises a tissue thickness that is in a particular range of tissue thicknesses, for example.

[0273] In certain instances, as illustrated in FIG. 89, the memory 4316 can store one or more predefined ranges of tissue thicknesses of tissue captured by the end effector 300; and predefined threshold forces associated with the predefined ranges of tissue thicknesses. In certain instances, each predefined threshold force may represent a minimum sharpness level of the cutting edge 182 that is suitable for transecting a captured tissue comprising a tissue thickness (Tx) encompassed by the range of tissue thicknesses that is associated with the predefined threshold force. In certain instances, if the force (Fx) required for the cutting edge 182 to transect the captured tissue, comprising the tissue thickness (Tx), exceeds the predefined threshold force associated with the predefined range of tissue thicknesses that encompasses the tissue thickness (Tx), the cutting edge 182 may not be sufficiently sharp to transect the captured tissue, for example.

[0274] In certain instances, the predefined threshold forces and their corresponding predefined ranges of tissue thicknesses can be stored in a database and/or a table on the memory 4316 such as, for example, a table 4342, as illustrated in FIG. 89. In certain instances, the processor 4314 can be configured to receive a measured value of the force (Fx) required for the cutting edge 182 to transect a captured tissue and a measured value of the tissue thickness (Tx) of the captured tissue. The processor 4314 may access the table 4342 to determine the predefined range of tissue thicknesses that encompasses the measured tissue thickness (Tx). In addition, the processor 4314 may compare the measured force (Fx) to the predefined threshold force associated with the predefined range of tissue thicknesses that encompasses the tissue thickness (Tx). In certain instances, if the measured force (Fx) exceeds the predefined threshold force, the processor 4314 may conclude that the cutting edge 182 may not be sufficiently sharp to transect the captured tissue, for example.

[0275] Further to the above, the processor 4314 (FIGS. 85, 86) may employ one or more tissue thickness sensing modules such as, for example, a tissue thickness sensing module 4336 to determine the thickness of the captured tissue. Various suitable tissue thickness sensing modules are described in the present disclosure. In addition, various tissue thickness sensing devices and methods, which are suitable for use with the present disclosure, are disclosed in U.S. Patent Application Publication No.

¹⁰ 2011/0155781, entitled SURGICAL CUTTING INSTRU-MENT THAT ANALYZES TISSUE THICKNESS, which is herein incorporated by reference in its entirety. [0276] In certain instances, the processor 4314 may employ the load cell 4334 to measure the force (Fx) re-

¹⁵ quired for the cutting edge 182 to transect a captured tissue comprising a tissue thickness (Tx). The reader will appreciate that that the force applied to the cutting edge 182 by the captured tissue, while the cutting edge 182 is engaged and/or in contact with the captured tissue, may

²⁰ increase as the cutting edge 182 is advanced against the captured tissue up to the force (Fx) at which the cutting edge 182 may transect the captured tissue. In certain instances, the processor 4314 may employ the load cell 4334 to continually monitor the force applied by the cap-

²⁵ tured tissue against the cutting edge 182 as the cutting edge 182 is advanced against the captured tissue. The processor 4314 may continually compare the monitored force to the predefined threshold force associated with the predefined tissue thickness range encompassing the

³⁰ tissue thickness (Tx) of the captured tissue. In certain instances, if the monitored force exceeds the predefined threshold force, the processor 4314 may conclude that the cutting edge is not sufficiently sharp to safely transect the captured tissue, for example.

³⁵ [0277] The method 4600 described in FIG. 88 outline various example actions that can be taken by the controller 4313 in the event it is determined that the cutting edge 182 is not be sufficiently sharp to safely transect the captured tissue, for example. In certain instances,

40 the controller 4312 may warn the user that the cutting edge 182 is too dull for safe use, for example, through the feedback system 1120, for example. In certain instances, the controller 4312 may employ the firing lockout mechanism 1122 to prevent advancement of the cutting

edge 182 upon concluding that the cutting edge 182 is not sufficiently sharp to safely transect the captured tissue, for example. In certain instances, the controller 4312 may employ the feedback system 1120 to provide instructions to the user for overriding the firing lockout
mechanism 1122, for example.

[0278] FIGS. 90, 91 illustrate various aspects of an apparatus, system, and method for employing a common controller with a plurality of motors in connection with a surgical instrument such as, for example, a motor-driven surgical instrument 4400. The surgical instrument 4400 is similar in many respects to other surgical instruments described by the present disclosure such as, for example, the surgical instrument 10 of FIG. 1 which is described

in greater detail above. The surgical instrument 4400 includes the housing 12, the handle assembly 14, the closure trigger 32, the interchangeable shaft assembly 200, and the end effector 300. Accordingly, for conciseness and clarity of disclosure, a detailed description of certain features of the surgical instrument 4400, which are common with the surgical instrument 10, will not be repeated here.

[0279] Referring still to FIGS. 90, 91, the surgical instrument 4400 may include a plurality of motors which can be activated to perform various functions in connection with the operation of the surgical instrument 4400. In certain instances, a first motor can be activated to perform a first function; a second motor can be activated to perform a second function; and a third motor can be activated to perform a third function. In certain instances, the plurality of motors of the surgical instrument 4400 can be individually activated to cause articulation, closure, and/or firing motions in the end effector 300 (FIGS. 1, 15). The articulation, closure, and/or firing motions can be transmitted to the end effector 300 through the interchangeable shaft assembly 200 (FIG. 1), for example. [0280] In certain instances, as illustrated in FIG. 91, the surgical instrument 4400 may include a firing motor 4402. The firing motor 4402 may be operably coupled to a firing drive assembly 4404 which can be configured to transmit firing motions generated by the firing motor 4402 to the end effector 300 (FIGS. 1, 14). In certain instances, the firing motions generated by the firing motor 4402 may cause the staples 191 to be deployed from the surgical staple cartridge 304 into tissue captured by the end effector 300 and/or the cutting edge 182 to be advanced to cut the captured tissue, for example.

[0281] In certain instances, as illustrated in FIG. 91, the surgical instrument 4400 may include an articulation motor 4406, for example. The articulation motor 4406 may be operably coupled to an articulation drive assembly 4408 which can be configured to transmit articulation motions generated by the articulation motor 4406 to the end effector 300 (FIGS. 1, 14). In certain instances, the articulation motions may cause the end effector 300 to articulate relative to the interchangeable shaft assembly 200 (FIG. 1), for example. In certain instances, the surgical instrument 4400 may include a closure motor, for example. The closure motor may be operably coupled to a closure drive assembly which can be configured to transmit closure motions to the end effector 300. In certain instances, the closure motions may cause the end effector 300 to transition from an open configuration to an approximated configuration to capture tissue, for example. The reader will appreciate that the motors described herein and their corresponding drive assemblies are intended as examples of the types of motors and/or driving assemblies that can be employed in connection with the present disclosure. The surgical instrument 4400 may include various other motors which can be utilized to perform various other functions in connection with the operation of the surgical instrument 4400.

[0282] As described above, the surgical instrument 4400 may include a plurality of motors which may be configured to perform various independent functions. In certain instances, the plurality of motors of the surgical instrument 4400 can be individually or separately activated to perform one or more functions while the other motors remain inactive. For example, the articulation motor 4406 can be activated to cause the end effector 300 (FIGS. 1, 14) to be articulated while the firing motor 4402

remains inactive. Alternatively, the firing motor 4402 can be activated to fire the plurality of staples 191 (FIG. 14) and/or advance the cutting edge 182 while the articulation motor 4406 remains inactive.

[0283] With reference to FIGS. 90, 91, in certain instances, the surgical instrument 4400 may include a common controller 4410 which can be employed with a plurality of motors 4402, 4406 of the surgical instrument 4400. In certain instances, the common controller 4410 may accommodate one of the plurality of motors at a

time. For example, the common controller 4410 can be separably couplable to the plurality of motors of the surgical instrument 4400 individually. In certain instances, a plurality of the motors of the surgical instrument 4400 may share one or more common controllers such as the

common controller 4410. In certain instances, a plurality of motors of the surgical instrument 4400 can be individually and selectively engaged the common controller 4410. In certain instances, the common controller 4410 can be selectively switched from interfacing with one of
 a plurality of motors of the surgical instrument 4400 to interfacing with another one of the plurality of motors of the surgical instrument 4400.

[0284] In at least one example, the common controller 4410 can be selectively switched between operable engagement with the articulation motor 4406 and operable engagement with the firing motor 4402. In at least one example, as illustrated in FIG. 90, a switch 4414 can be moved or transitioned between a plurality of positions and/or states such as a first position 4416 and a second position 4418, for example. In the first position 4416, the switch 4414 may electrically couple the common controller 4410 to the articulation motor 4406; and in the second position 4418, the switch 4414 may electrically couple the common controller 4410 to the articulation motor 4402, for

45 example. In certain instances, the common controller 4410 can be electrically coupled to the articulation motor 4406, while the switch 4414 is in the first position 4416, to control the operation of the articulation motor 4406 to articulate the end effector 300 (FIGS. 1, 15) to a desired 50 position. In certain instances, the common controller 4410 can be electrically coupled to the firing motor 4402, while the switch 4414 is in the second position 4418, to control the operation of the firing motor 4402 to fire the plurality of staples 191 (FIG. 14) and/or advance the cut-55 ting edge 182 (FIG. 14), for example. In certain instances, the switch 4414 may be a mechanical switch, an electromechanical switch, a solid state switch, or any suitable switching mechanism.

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[0285] Referring now to FIG. 91, an outer casing of the handle assembly 14 of the surgical instrument 4400 is removed and several features and elements of the surgical instrument 4400 are also removed for clarity of disclosure. In certain instances, as illustrated in FIG. 91, the surgical instrument 4400 may include an interface 4412 which can be selectively transitioned between a plurality of positions and/or states. In a first position and/or state, the interface 4412 may couple the common controller 4410 (FIG. 90) to a first motor such as, for example, the articulation motor 4406; and in a second position and/or state, the interface 4412 may couple the common controller 4410 to a second motor such as, for example, the firing motor 4402. Additional positions and/or states of the interface 4412 are contemplated by the present disclosure.

[0286] In certain instances, the interface 4412 is movable between a first position and a second position, wherein the common controller 4410 (FIG. 90) is coupled to a first motor in the first position and a second motor in the second position. In certain instances, the common controller 4410 is decoupled from first motor as the interface 4412 is moved from the first position; and the common controller 4410 is decoupled from second motor as the interface 4412 is moved from the second position. In certain instances, a switch or a trigger can be configured to transition the interface 4412 between the plurality of positions and/or states. In certain instances, a trigger can be movable to simultaneously effectuate the end effector and transition the common controller 4410 from operable engagement with one of the motors of the surgical instrument 4400 to operable engagement with another one of the motors of the surgical instrument 4400. [0287] In at least one example, as illustrated in FIG. 91, the closure trigger 32 can be operably coupled to the interface 4412 and can be configured to transition the interface 4412 between a plurality of positions and/or states. As illustrated in FIG. 91, the closure trigger 32 can be movable, for example during a closure stroke, to transition the interface 4412 from a first position and/or state to a second position and/or state while transitioning the end effector 300 to an approximated configuration to capture tissue by the end effector, for example.

[0288] In certain instances, in the first position and/or state, the common controller 4410 can be electrically coupled to a first motor such as, for example, the articulation motor 4406, and in the second position and/or state, the common controller 4410 can be electrically coupled to a second motor such as, for example, the firing motor 4402. In the first position and/or state, the common controller 4410 may be engaged with the articulation motor 4406 to allow the user to articulate the end effector 300 (FIGS. 1, 15) to a desired position; and the common controller 4410 may remain engaged with the articulation motor 4406 until the closure trigger 32 is actuated. As the user actuates the closure trigger 32 to capture tissue by the end effector 300 at the desired position, the interface 4412 can be transitioned or shifted

to transition the common controller 4410 from operable engagement with the articulation motor 4406, for example, to operable engagement with the firing motor 4402, for example. Once operable engagement with the firing motor 4402 is established, the common controller 4410 may take control of the firing motor 4402; and the common controller 4410 may activate the firing motor 4402, in response to user input, to fire the plurality of staples 191 (FIG. 14) and/or advance the cutting edge 182 (FIG. 14), for example.

[0289] In certain instances, as illustrated in FIG. 91, the common controller 4410 may include a plurality of electrical and/or mechanical contacts 4411 adapted for coupling engagement with the interface 4412. The plu-

¹⁵ rality of motors of the surgical instrument 4400, which share the common controller 4410, may each comprise one or more corresponding electrical and/or mechanical contacts 4413 adapted for coupling engagement with the interface 4412, for example.

20 [0290] In various instances, the motors of the surgical instrument 4400 can be electrical motors. In certain instances, one or more of the motors of the surgical instrument 4400 can be a DC brushed driving motor having a maximum rotation of, approximately, 25,000 RPM, for

25 example. In other arrangements, the motors of the surgical instrument 4400 may include one or more motors selected from a group of motors comprising a brushless motor, a cordless motor, a synchronous motor, a stepper motor, or any other suitable electric motor.

30 [0291] In various instances, as illustrated in FIG. 90, the common controller 4410 may comprise a motor driver 4426 which may comprise one or more H-Bridge field-effect transistors (FETs). The motor driver 4426 may modulate the power transmitted from a power source
 35 4428 to a motor coupled to the common controller 4410 based on input from a controller 4420 ("microcontroller"), for example. In certain instances, the controller 4420 can be employed to determine the current drawn by the motor, for example, while the motor is coupled to the common controller 4410, as described above.

[0292] In certain instances, the controller 4420 may include a processor 4422 ("microprocessor") and one or more computer readable mediums or memory 4424 units ("memory"). In certain instances, the memory 4424 may

⁴⁵ store various program instructions, which when executed may cause the processor 4422 to perform a plurality of functions and/or calculations described herein. In certain instances, one or more of the memory 4424 may be coupled to the processor 4422, for example.

⁵⁰ [0293] In certain instances, the power source 4428 can be employed to supply power to the controller 4420, for example. In certain instances, the power source 4428 may comprise a battery (or "battery pack" or "power pack"), such as a Li ion battery, for example. In certain ⁵⁵ instances, the battery pack may be configured to be releasably mounted to the handle assembly 14 for supplying power to the surgical instrument 4400. A number of battery cells connected in series may be used as the

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[0294] In various instances, the processor 4422 may control the motor driver 4426 to control the position, direction of rotation, and/or velocity of a motor that is coupled to the common controller 4410. In certain instances, the processor 4422 can signal the motor driver 4426 to stop and/or disable a motor that is coupled to the common controller 4410. It should be understood that the term processor as used herein includes any suitable processor, controller, or other basic computing device that incorporates the functions of a computer's central processing unit (CPU) on an integrated circuit or at most a few integrated circuits. The processor is a multipurpose, programmable device that accepts digital data as input, processes it according to instructions stored in its memory, and provides results as output. It is an example of sequential digital logic, as it has internal memory. Processors operate on numbers and symbols represented in the binary numeral system. In one instance, the processor 4422 may be a single core or multicore controller LM4F230H5QR as described in connection with FIGS. 15-17B.

[0295] In certain instances, the memory 4424 may include program instructions for controlling each of the motors of the surgical instrument 4400 that are couplable to the common controller 4410. For example, the memory 4424 may include program instructions for controlling the articulation motor 4406. Such program instructions may cause the processor 4422 to control the articulation motor 4406 to articulate the end effector 300 in accordance with user input while the articulation motor 4406 is coupled to the common controller 4410. In another example, the memory 4424 may include program instructions for controlling the firing motor 4402. Such program instructions may cause the processor 4422 to control the firing motor 4402 to fire the plurality of staples 191 and/or advance the cutting edge 182 in accordance with user input while the firing motor 4402 is coupled to the common controller 4410.

[0296] In certain instances, one or more mechanisms and/or sensors such as, for example, sensors 4430 can be employed to alert the processor 4422 to the program instructions that should be used in a particular setting. For example, the sensors 4430 may alert the processor 4422 to use the program instructions associated with articulation of the end effector 300 (FIGS. 1, 14) while the common controller 4410 is coupled to the articulation motor 4406; and the sensors 4430 may alert the processor 4422 to use the program instructions associated with firing the surgical instrument 4400 while the common controller 4410 is coupled to the firing motor 4402. In certain instances, the sensors 4430 may comprise position sensors which can be employed to sense the position of the switch 4414, for example. Accordingly, the processor 4422 may use the program instructions associated with articulation of the end effector 300 upon detecting,

through the sensors 4430 for example, that the switch 4414 is in the first position 4416; and the processor 4422 may use the program instructions associated with firing the surgical instrument 4400 upon detecting, through the sensors 4430 for example, that the switch 4414 is in the

second position 4418. [0297] Referring now to FIG. 92, an outer casing of the

surgical instrument 4400 is removed and several features and elements of the surgical instrument 4400 are also removed for clarity of disclosure. As illustrated in FIG.

92, the surgical instrument 4400 may include a plurality of sensors which can be employed to perform various functions in connection with the operation of the surgical instrument 4400. For example, as illustrated in FIG. 92,

¹⁵ the surgical instrument 4400 may include sensors A, B, and/or C. In certain instances, the sensor A can be employed to perform a first function, for example; the sensor B can be employed to perform a second function, for example; and the sensor C can be employed to perform

²⁰ a third function, for example. In certain instances, the sensor A can be employed to sense a thickness of the tissue captured by the end effector 300 (FIGS. 1, 14) during a first segment of a closure stroke; the sensor B can be employed to sense the tissue thickness during a

second segment of the closure stroke following the first segment; and the sensor C can be employed to sense the tissue thickness during a third segment of the closure stroke following the second segment, for example. In certain instances, the sensors A, B, and C can be disposed along the end effector 300, for example.

[0298] In certain instances, the sensors A, B, and C can be arranged, as illustrated in FIG. 94, such that the sensor A is disposed proximal to the sensor B, and the sensor C is disposed proximal to the sensor B, for exam-

ple. In certain instances, as illustrated in FIG. 92, the sensor A can sense the tissue thickness of the tissue captured by the end effector 300 at a first position; the sensor B can sense the tissue thickness of the tissue captured by the end effector 300 at a second position
distal to the first position; and the sensor C can sense the tissue thickness of the tissue captured by the end effector 300 at a third position distal to the second posi-

tion, for example. The reader will appreciate that the sensors described herein are intended as examples of the
 types of sensors which can be employed in connection with the present disclosure. Other suitable sensors and sensing arrangements can be employed by the present disclosure.

[0299] In certain instances, the surgical instrument
⁵⁰ 4400 may include a controller 4450 which can be similar in many respects to the common controller 4410. For example, the controller 4450, like the common controller 4410, may comprise the controller 4420, the processor 4422, and/or the memory 4424. In certain instances, the
⁵⁵ power source 4428 can supply power to the controller 4450, for example. In certain instances, the surgical instrument 4400 may include a plurality of sensors such as the sensors A, B, and C, for example, which can ac-

tivated to perform various functions in connection with the operation of the surgical instrument 4400. In certain instances, one of the sensors A, B, and C, for example, can be individually or separately activated to perform one or more functions while the other sensors remain inactive. In certain instances, a plurality of sensors of the surgical instrument 4400 such as, for example, the sensors A, B, and C may share the controller 4450. In certain instances, only one of the sensors A, B, and C can be coupled to the controller 4450 at a time. In certain instances, the plurality of sensors of the surgical instrument 4400 can be individually and separately couplable to the controller 4450, for example. In at least one example, the controller 4450 can be selectively switched between operable engagement with sensor A, Sensor B, and/or Sensor C.

[0300] In certain instances, as illustrated in FIG. 92, the controller 4450 can be disposed in the handle assembly 14, for example, and the sensors that share the controller 4450 can be disposed in the end effector 300 (FIGS. 1, 14), for example. The reader will appreciate that the controller 4450 and/or the sensors that share the controller 4450 are not limited to the above identified positions. In certain instances, the controller 4450 and the sensors that share the controller 300, for example. Other arrangements for the positions of the controller 4450 and/or the sensors that share the controller 4450 and/or the sensors that share the controller 4450 and the sensors disposed in the end effector 300, for example. Other arrangements for the positions of the controller 4450 and/or the sensors that share the controller 4450 and the sensors that share the controller 4450 and/or the sensors that share the controller 4450 are contemplated by the present disclosure.

[0301] In certain instances, as illustrated in FIG. 92, an interface 4452 can be employed to manage the coupling and/or decoupling of the sensors of the surgical instrument 4400 to the controller 4450. In certain instances, the interface 4452 can be selectively transitioned between a plurality of positions and/or states. In a first position and/or state, the interface 4452 may couple the controller 4450 to the sensor A, for example; in a second position and/or state, the interface 4452 may couple the controller 4450 to the sensor B, for example; and in a third position and/or state, the interface 4452 may couple the controller 4450 to the sensor C, for example. Additional positions and/or states of the interface 4452 are contemplated by the present disclosure.

[0302] In certain instances, the interface 4452 is movable between a first position, a second position, and/or a third position, for example, wherein the controller 4450 is coupled to a first sensor in the first position, a second sensor in the second position, and a third sensor in the third position. In certain instances, the controller 4450 is decoupled from first sensor as the interface 4452 is moved from the first position; the controller 4450 is decoupled from second sensor as the interface 4452 is moved from the second position; and the controller 4450 is decoupled from third sensor as the interface 4452 is moved from the third position. In certain instances, a switch or a trigger can be configured to transition the interface 4452 between the plurality of positions and/or states. In certain instances, a trigger can be movable to simultaneously effectuate the end effector and transition the controller 4450 from operable engagement with one of the sensors that share the controller 4450 to operable engagement with another one of the sensors that share the controller 4450, for example.

[0303] In at least one example, as illustrated in FIG. 92, the closure trigger 32 can be operably coupled to the interface 4452 and can be configured to transition the interface 4452 between a plurality of positions and/or

¹⁰ states. As illustrated in FIG. 92, the closure trigger 32 can be moveable between a plurality of positions, for example during a closure stroke, to transition the interface 4452 between a first position and/or state wherein the controller 4450 is electrically coupled to the sensor A, for

¹⁵ example, a second position and/or state wherein the controller 4450 is electrically coupled to the sensor B, for example, and/or a third position and/or state wherein the controller 4450 is electrically coupled to the sensor C, for example.

20 [0304] In certain instances, a user may actuate the closure trigger 32 to capture tissue by the end effector 300. Actuation of the closure trigger may cause the interface 4452 to be transitioned or shifted to transition the controller 4450 from operable engagement with the sensor

A, for example, to operable engagement with the sensor B, for example, and/or from operable engagement with sensor B, for example, to operable engagement with sensor C, for example.

[0305] In certain instances, the controller 4450 may be
 coupled to the sensor A while the closure trigger 32 is in a first actuated position. As the closure trigger 32 is actuated past the first actuated position and toward a second actuated position, the controller 4450 may be decoupled from the sensor A. Alternatively, the controller 4450

³⁵ may be coupled to the sensor A while the closure trigger 32 is in an unactuated position. As the closure trigger 32 is actuated past the unactuated position and toward a second actuated position, the controller 4450 may be decoupled from the sensor A. In certain instances, the con-

40 troller 4450 may be coupled to the sensor B while the closure trigger 32 is in the second actuated position. As the closure trigger 32 is actuated past the second actuated position and toward a third actuated position, the controller 4450 may be decoupled from the sensor B. In

⁴⁵ certain instances, the controller 4450 may be coupled to the sensor C while the closure trigger 32 is in the third actuated position.

[0306] In certain instances, as illustrated in FIG. 92, the controller 4450 may include a plurality of electrical and/or mechanical contacts 4451 adapted for coupling engagement with the interface 4452. The plurality of sensors of the surgical instrument 4400, which share the controller 4450, may each comprise one or more corresponding electrical and/or mechanical contacts 4453
⁵⁵ adapted for coupling engagement with the interface 4452, for example.

[0307] In certain instances, the processor 4422 may receive input from the plurality of sensors that share the

controller 4450 while the sensors are coupled to the interface 4452. For example, the processor 4422 may receive input from the sensor A while the sensor A is coupled to the controller 4450; the processor 4422 may receive input from the sensor B while the sensor B is coupled to the controller 4450; and the processor 4422 may receive input from the sensor C while the sensor C is coupled to the controller 4450. In certain instances, the input can be a measurement value such as, for example, a measurement value of a tissue thickness of tissue captured by the end effector 300 (FIGS. 1, 15). In certain instances, the processor 4422 may store the input from one or more of the sensors A, B, and C on the memory 4424. In certain instances, the processor 4422 may perform various calculations based on the input provided by the sensors A, B, and C, for example.

[0308] FIGS. 93A and 93B illustrate one aspect of an end effector 5300 comprising a staple cartridge 5306 that further comprises two light-emitting diodes 5310 (LEDs). FIG. 93A illustrates an end effector 5300 comprising one LED 5310 located on either side of the cartridge deck 5308. FIG. 91B illustrates a three-quarter angle view of the end effector 5300 with the anvil 5302 in an open position, and one LED 5310 located on either side of the cartridge deck 5308. The end effector 5300 is similar to the end effector 300 (FIGS. 1, 15) described above. The end effector comprises an anvil 5302, pivotally coupled to a jaw member or elongated channel 5304. The elongated channel 5304 is configured to receive the staple cartridge 5306 therein. The staple cartridge 5306 comprises a plurality of staples (not shown). The plurality of staples are deployable from the staple cartridge 5306 during a surgical operation. The staple cartridge 5306 further comprises two LEDs 5310 mounted on the upper surface, or cartridge deck 5308 of the staple cartridge 5306. The LEDs 5310 are mounted such that they will be visible when the anvil 5302 is in a closed position. Furthermore, the LEDs 5310 can be sufficiently bright to be visible through any tissue that may be obscuring a direct view of the LEDs 5310. Additionally, one LED 5310 can be mounted on either side of the staple cartridge 5306 such that at least one LED 5310 is visible from either side of the end effector 5300. The LED 5310 can be mounted near the proximal end of the staple cartridge 530, as illustrated, or may be mounted at the distal end of the staple cartridge 5306.

[0309] The LEDs 5310 may be in communication with a processor or controller, such as, for instance, controller 1500 (FIG. 19). The controller 1500 can be configured to detect a property of tissue compressed by the anvil 5302 against the cartridge deck 5308. Tissue that is enclosed by the end effector 5300 may change height as fluid within the tissue is exuded from the tissue's layers. Stapling the tissue before it has sufficiently stabilized may affect the effectiveness of the staples. Tissue stabilization is typically communicates as a rate of change, where the rate of change indicates how rapidly the tissue enclosed by the end effector is changing height. **[0310]** The LEDs 5310 mounted to the staple cartridge 5306, in the view of the operator of the instrument, can be used to indicate rate at which the enclosed tissue is stabilizing and/or whether the tissue has reached a stable state. The LEDs 5310 can, for example, be configured to flash at a rate that directly correlates to the rate of

stabilization of the tissue, that is, can flash quickly initially, flash slower as the tissue stabilizes, and remain steady when the tissue is stable. Alternatively, the LEDs 5310
can flash slowly initially, flash more quickly as the tissue

stabilizes, and turn off when the tissue is stable.
[0311] The LEDs 5310 mounted on the staple cartridge 5306 can be used additionally or optionally to indicate other information. Examples of other information include,

¹⁵ but are not limited to: whether the end effector 5300 is enclosing a sufficient amount of tissue, whether the staple cartridge 5306 is appropriate for the enclosed tissue, whether there is more tissue enclosed than is appropriate for the staple cartridge 5306, whether the staple cartridge

²⁰ 5306 is not compatible with the surgical instrument, or any other indicator that would be useful to the operator of the instrument. The LEDs 5310 can indicate information by either flashing at a particular rate, turning on or off at a particular instance, lighting in different colors for

different information. The LEDs 5310 can alternatively or additionally be used to illuminate the area of operation. In some aspects the LEDs 5310 can be selected to emit ultraviolet or infrared light to illuminate information not visible under normal light, where that information is print ed on the staple cartridge located in the end effector 5300 or on a tissue compensator (not illustrated). Alternatively or additionally, the staples can be coated with a fluorescing dye and the wavelength of the LEDs 5310 chosen so

that the LEDs 5310 cause the fluorescing dye to glow.
By illuminating the staples with the LEDs 5310 allows the operator of the instrument to see the staples after they have been driven.

[0312] FIGS. 94A and 94B illustrate one aspect of the end effector 5300 comprising a staple cartridge 5356 that
⁴⁰ further comprises a plurality of LEDs 5360. FIG. 92A illustrates a side angle view of the end effector 5300 with the anvil 5302 in a closed position. The illustrated aspect comprises, by way of example, a plurality of LEDs 5360 located on either side of the cartridge deck 5358. FIG.

45 92B illustrates a three-quarter angle view of the end effector 5300 with the anvil 5302 in an open position, illustrating a plurality of LEDs 5360 located on either side of the cartridge deck 5358. The staple cartridge 5356 comprises a plurality of LEDs 5360 mounted on the cartridge 50 deck 5358 of the staple cartridge 5356. The LEDs 5360 are mounted such that they will be visible when the anvil 5302 is in a closed position. Furthermore, the LEDs6 530 can be sufficiently bright to be visible through any tissue that may be obscuring a direct view of the LEDs 5360. 55 Additionally, the same number of LEDs 5360 can be mounted on either side of the staple cartridge 5356 such that the same number of LEDs 5360 is visible from either side of the end effector 5300. The LEDs 5360 can be

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mounted near the proximal end of the staple cartridge 5356, as illustrated, or may be mounted at the distal end of the staple cartridge 5356.

[0313] The LEDs 5360 may be in communication with a processor or controller, such as, for instance, controller 1500 of FIG. 15. The controller 1500 can be configured to detect a property of tissue compressed by the anvil 5302 against the cartridge deck 5358, such as the rate of stabilization of the tissue, as described above. The LEDs 5360 can be used to indicate the rate at which the enclose tissue is stabilizing and/or whether the tissue has reached a stable state. The LEDs 5360 can be configured, for instance, to light in sequence starting at the proximal end of the staple cartridge 5356 with each subsequent LED 5360 lighting at the rate at which the enclosed tissue is stabilizing; when the tissue is stable, all the LEDs 5360 can be lit. Alternatively, the LEDs 5360 can light in sequence beginning at the distal end of the staple cartridge 5356. Yet another alternative is for the LEDs 5360 to light in a sequential, repeating sequence, with the sequence starting at either the proximal or distal end of the LEDs 5360. The rate at which the LEDs 5360 light and/or the speed of the repeat can indicate the rate at which the enclosed tissue is stabilizing. It is understood that these are only examples of how the LEDs 5360 can indicate information about the tissue, and that other combinations of the sequence in which the LEDs 5360 light, the rate at which they light, and or their on or off state are possible. It is also understood that the LEDs 5360 can be used to communicate some other information to the operator of the surgical instrument, or to light the work area, as described above.

[0314] FIGS. 95A and 95B illustrate one aspect of the end effector 5300 comprising a staple cartridge 5406 that further comprises a plurality of LEDs 5410. FIG. 93A illustrates a side angle view of the end effector 5300 with the anvil 5302 in a closed position. The illustrated aspect comprises, by way of example, a plurality of LEDs 5410 from the proximal to the distal end of the staple cartridge 5406, on either side of the cartridge deck 5408. FIG. 93B illustrates a three-quarter angle view of the end effector 5300 with the anvil 5302 in an open position, illustrating a plurality of LEDs 5410 from the proximal to the distal end of the staple cartridge 5406, and on either side of the cartridge deck 5408. The staple cartridge 5406 comprises a plurality of LEDs 5410 mounted on the cartridge deck 5408 of the staple cartridge 5406, with the LEDs 5410 placed continuously from the proximal to the distal end of the staple cartridge 5406. The LEDs 5410 are mounted such that they will be visible when the anvil 5302 is in a closed position. The same number of LEDs 5410 can be mounted on either side of the staple cartridge 5406 such that the same number of LEDs 5410 is visible from either side of the end effector 5300.

[0315] The LEDs 5410 can be in communication with a processor or controller, such as, for instance, controller 1500 of FIG. 15. The controller 1500 can be configured to detect a property of tissue compressed by the anvil

5302 against the cartridge deck 5408, such as the rate of stabilization of the tissue, as described above. The LEDs 5410 can be configured to be turned on or off in sequences or groups as desired to indicate the rate of stabilization of the tissue and/or that the tissue is stable. The LEDs 5410 can further be configured communicate some other information to the operator of the surgical

instrument, or to light the work area, as described above. Additionally or alternatively, the LEDs 5410 can be configured to indicate which areas of the end effector 5300 contain stable tissue, and or what areas of the end ef-

fector 5300 are enclosing tissue, and/or if those areas are enclosing sufficient tissue. The LEDs 5410 can further be configured to indicate if any portion of the en-

¹⁵ closed tissue is unsuitable for the staple cartridge 5406. [0316] Referring now primarily to FIGS. 96 and 97, the power assembly 2096 may include a power modulator control 2106 which may comprise, for example, one or more field-effect transistors (FETs), a Darlington array,

an adjustable amplifier, and/or any other power modulator. The power assembly controller 2100 may actuate the power modulator control 2106 to set the power output of the battery 2098 to the power requirement of the inter-changeable working assembly 2094 in response to the
 signal generated by working assembly controller 2102

while the interchangeable working assembly 2094 is coupled to the power assembly 2096.

[0317] Still referring primarily to FIGS. 96 and 97, the power assembly controller 2100 can be configured to monitor power transmission from the power assembly 2096 to the interchangeable working assembly 2094 for the one or more signals generated by the working assembly controller 2102 of the interchangeable working assembly 2094 while he interchangeable working assembly 2094 is coupled to the power assembly 2096. As

illustrated in FIG. 96, the power assembly controller 2100 may utilize a voltage monitoring mechanism for monitoring the voltage across the battery 2098 to detect the one or more signals generated by the working assembly con-

40 troller 2102, for example. In certain instances, a voltage conditioner can be utilized to scale the voltage of the battery 2098 to be readable by an Analog to Digital Converter (ADC) of the power assembly controller 2100. As illustrated in FIG. 96, the voltage conditioner may com-

⁴⁵ prise a voltage divider 2108 which can create a reference voltage or a low voltage signal proportional to the voltage of the battery 2098 which can be measured and reported to the power assembly controller 2100 through the ADC, for example.

50 [0318] In other circumstances, as illustrated in FIG. 97, the power assembly 2096 may comprise a current monitoring mechanism for monitoring current transmitted to the interchangeable working assembly 2094 to detect the one or more signals generated by the working assembly controller 2102, for example. In certain instances, the power assembly 2096 may comprise a current sensor 2110 which can be utilized to monitor current transmitted to the interchangeable working assembly 2094. The

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monitored current can be reported to the power assembly controller 2100 through an ADC, for example. In other circumstances, the power assembly controller 2100 may be configured to simultaneously monitor both of the current transmitted to the interchangeable working assembly 2094 and the corresponding voltage across the battery 2098 to detect the one or more signals generated by the working assembly controller 2102. The reader will appreciate that various other mechanisms for monitoring current and/or voltage can be utilized by the power assembly controller 2100 to detect the one or more signals generated by the working assembly controller 2102; all such mechanisms are contemplated by the present disclosure.

[0319] Referring to FIG. 98, the controller 13002 may generally comprise a processor 13008 ("microprocessor") and one or more memory units 13010 operationally coupled to the processor 13008. By executing instruction code stored in the memory 13010, the processor 13008 may control various components of the surgical instrument 12200, such as the motor 12216, various drive systems, and/or a user display, for example. The controller 13002 may be implemented using integrated and/or discrete hardware elements, software elements, and/or a combination of both. Examples of integrated hardware elements may include processors, microprocessors, controllers, integrated circuits, application specific integrated circuits (ASIC), programmable logic devices (PLD), digital signal processors (DSP), field programmable gate arrays (FPGA), logic gates, registers, semiconductor devices, chips, microchips, chip sets, controllers, system-on-chip (SoC), and/or system-in-package (SIP). Examples of discrete hardware elements may include circuits and/or circuit elements such as logic gates, field effect transistors, bipolar transistors, resistors, capacitors, inductors, and/or relays. In certain instances, the controller 13002 may include a hybrid circuit comprising discrete and integrated circuit elements or components on one or more substrates, for example. In certain instances, the controller 13002 may be a single core or multicore controller LM4F230H5QR as described in connection with FIGS. 15-17B.

[0320] In various forms, the motor 12216 may be a DC brushed driving motor having a maximum rotation of, approximately, 25,000 RPM, for example. In other arrangements, the motor 12216 may include a brushless motor, a cordless motor, a synchronous motor, a stepper motor, or any other suitable electric motor. A battery 12218 (or "power source" or "power pack"), such as a Li ion battery, for example, may be coupled to the housing 12212 to supply power to the motor 12216, for example.

[0321] Referring again to FIG. 98, the surgical instrument 12200 may include a motor controller 13005 in operable communication with the controller 13002. The motor controller 13005 can be configured to control a direction of rotation of the motor 12216. In certain instances, the motor controller 13005 may be configured to determine the voltage polarity applied to the motor 12216 by

the battery 12218 and, in turn, determine the direction of rotation of the motor 12216 based on input from the controller 13002. For example, the motor 12216 may reverse the direction of its rotation from a clockwise direction to

⁵ a counterclockwise direction when the voltage polarity applied to the motor 12216 by the battery 12218 is reversed by the motor controller 13005 based on input from the controller 13002. In addition, the motor 12216 can be operably coupled to an articulation drive which can be

¹⁰ driven by the motor 12216 distally or proximally depending on the direction in which the motor 12216 rotates, for example. Furthermore, the articulation drive can be operably coupled to the end effector 12208 such that, for example, the axial translation of the articulation drive

¹⁵ proximally may cause the end effector 12208 to be articulated in the counterclockwise direction, for example, and/or the axial translation of the articulation drive distally may cause the end effector 12208 to be articulated in the clockwise direction, for example.

20 [0322] In the aspect illustrated in FIG. 99, an interface 3001 comprises multiple switches 3004A-C, 3084B wherein each of the switches 3004A-C is coupled to the controller 3002 via one of three electrical circuits 3006A-C, respectively, and switch 3084B is coupled to the con-

²⁵ troller 3002 via circuit 3084A. The reader will appreciate that other combinations of switches and circuits can be utilized with the interface 3001.

[0323] Further to the above, the controller 3002 may comprise a processor 3008 and/or one or more memory 3010 units. By executing instruction code stored in the memory 3010, the processor 3008 may control various components of the surgical instrument, such as the electric motor 1102 and/or a user display. The controller 3002 may be implemented using integrated and/or discrete hardware elements, software elements, and/or a combi-

nation of both. Examples of integrated hardware elements may include processors, microprocessors, controllers, integrated circuits, application specific integrated circuits (ASIC), programmable logic devices (PLD), dig-

40 ital signal processors (DSP), field programmable gate arrays (FPGA), logic gates, registers, semiconductor devices, chips, microchips, chip sets, controller, systemon-chip (SoC), and/or system-in-package (SIP). Examples of discrete hardware elements may include circuits

and/or circuit elements (e.g., logic gates, field effect transistors, bipolar transistors, resistors, capacitors, inductors, relay and so forth). In other aspects, the controller 3002 may include a hybrid circuit comprising discrete and integrated circuit elements or components on one or
 more substrates, for example.

[0324] Referring again to FIG. 99, the surgical instrument 1010 may include a motor controller 3005 in operable communication with the controller 3002. The motor controller 3005 can be configured to control a direction of rotation of the electric motor 1102. For example, the electric motor 1102 can be powered by a battery such as, for example, the battery 1104 and the controller 3002 may be configured to determine the voltage polarity ap-

plied to the electric motor 1102 by the battery 1104 and, in turn, the direction of rotation of the electric motor 1102 based on input from the controller 3002. For example, the electric motor 1102 may reverse the direction of its rotation from a clockwise direction to a counterclockwise direction when the voltage polarity applied to the electric motor 1102 by the battery 1104 is reversed by the motor controller 3005 based on input from the controller 3002. Examples of suitable motor controllers are described elsewhere in this document and include but are not limited to the driver 7010 (FIG. 100).

[0325] In addition, as described elsewhere in this document in greater detail, the electric motor 1102 can be operably coupled to an articulation drive. In use, the electric motor 1102 can drive the proximal articulation drive distally or proximally depending on the direction in which the electric motor 1102 rotates. Furthermore, the proximal articulation drive can be operably coupled to the end effector 1300 such that, for example, the axial translation of the proximal articulation drive 10030 proximally may cause the end effector 1300 to be articulated in the counterclockwise direction, for example, and/or the axial translation of the proximal articulation drive 10030 distally may cause the end effector 1300 to be articulated in the clockwise direction, for example.

[0326] Further to the above, referring again to FIG. 99, the interface 3001 can be configured such that the switch 3004A can be dedicated to clockwise articulation of the end effector 1300 and the switch 3004B can be dedicated to counterclockwise articulation of the end effector 1300. For example, the operator may articulate the end effector 1300 in the clockwise direction by closing the switch 3004A which may signal the controller 3002 to cause the electric motor 1102 to rotate in the clockwise direction thereby, as a result, causing the proximal articulation drive 10030 to be advanced distally and causing the end effector 1300 to be articulated in the clockwise direction. In another example, the operator may articulate the end effector 1300 in the counterclockwise direction by closing the switch 3004B which may signal the controller 3002 to cause the electric motor 1102 to rotate in the counterclockwise direction, for example, and retracting the proximal articulation drive 10030 proximally to articulate the end effector 1300 to in the counterclockwise direction.

[0327] As shown in FIG. 100, a sensor arrangement 7002 provides a unique position signal corresponding to the location of the longitudinally-movable drive member 1111. The electric motor 1102 can include a rotatable shaft 7016 that operably interfaces with a gear assembly 7014 that is mounted in meshing engagement with a with a set, or rack, of drive teeth on the longitudinally-movable drive member 1111. With reference also to FIG. 101, the sensor element 7026 may be operably coupled to the gear assembly 7106 such that a single revolution of the sensor element 7026 corresponds to some linear longitudinal translation of the longitudinally-movable drive member 1111, as described in more detail hereinbelow. In one aspect, an arrangement of gearing and sensors

can be connected to the linear actuator via a rack and pinion arrangement, or a rotary actuator via a spur gear or other connection. For aspects comprising a rotary screw-drive configuration where a larger number of turns would be required, a high reduction gearing arrangement

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between the drive member and the sensor, like a worm and wheel, may be employed. [0328] In accordance one aspect of the present disclo-

sure, the sensor arrangement 7002 for the absolute positioning system 7000 provides a position sensor 7012 that is more robust for use with surgical devices. By providing a unique position signal or value for each possible actuator position, such arrangement eliminates the need for a zeroing or calibration step and reduces the possi-

bility of negative design impact in the cases where noise or power brown-out conditions may create position sense errors as in conventional rotary encoder configurations.
 [0329] In one aspect, the sensor arrangement 7002 for the absolute positioning system 7000 replaces conven-

tional rotary encoders typically attached to the motor rotor and replaces it with a position sensor 7012 which generates a unique position signal for each rotational position in a single revolution of a sensor element associated with the position sensor 7012. Thus, a single revolution of a sensor element associated with the position sensor 7012

is equivalent to a longitudinal linear displacement d1 of the of the longitudinally-movable drive member 1111. In other words, d1 is the longitudinal linear distance that the longitudinally-movable drive member 1111 moves from
30 point "a" to point "b" after a single revolution of a sensor

element coupled to the longitudinally-movable drive member 1111. The sensor arrangement 7002 may be connected via a gear reduction that results in the position sensor 7012 completing only a single turn for the full stroke of the longitudinally-movable drive member 1111. With a suitable gear ratio, the full stroke of the longitudi-

nally-movable drive member 1111 can be represented in one revolution of the position sensor 7012.

[0330] A series of switches 7022a to 7022n, where n
 is an integer greater than one, may be employed alone or in combination with gear reduction to provide a unique position signal for more than one revolution of the position sensor 7012. The state of the switches 7022a-7022n are fed back to a controller 7004 which applies logic to de-

⁴⁵ termine a unique position signal corresponding to the longitudinal linear displacement d1 + d2 + ... dn of the longitudinally-movable drive member 1111.

[0331] Accordingly, the absolute positioning system 7000 provides an absolute position of the longitudinally⁵⁰ movable drive member 1111 upon power up of the instrument without retracting or advancing the longitudinally-movable drive member 1111 to a reset (zero or home) position as may be required with conventional rotary encoders that merely count the number of steps forwards or backwards that motor has taken to infer the position of a device actuator, drive bar, knife, and the like.
[0332] In various aspects, the position sensor 7012 of the sensor arrangement 7002 may comprise one or more

magnetic sensor, analog rotary sensor like a potentiometer, array of analog Hall-effect elements, which output a unique combination of position signals or values, among others, for example.

[0333] In various aspects, the controller 7004 may be programmed to perform various functions such as precise control over the speed and position of the knife and articulation systems. Using the known physical properties, the controller 7004 can be designed to simulate the response of the actual system in the software of the controller 7004. The simulated response is compared to (noisy and discrete) measured response, which is used for actual feedback decisions. The observed response is a favorable, tuned, value that balances the smooth, continuous nature of the simulated response with the measured response, which can detect outside influences on the system.

[0334] In various aspects, the absolute positioning system 7000 may further comprise and/or be programmed to implement the following functionalities. A feedback controller, which can be one of any feedback controllers, including, but not limited to: PID, state feedback and adaptive. A power source converts the signal from the feedback controller into a physical input to the system, in this case voltage. Other examples include, but are not limited to pulse width modulated (PWMed) voltage, current and force. The electric motor 1102 may be a brushed DC motor with a gearbox and mechanical links to an articulation or knife system. Other sensor(s) 7018 may be provided to measure physical parameters of the physical system in addition to position measured by the position sensor 7012. Since it is a digital signal (or connected to a digital data acquisition system) its output will have finite resolution and sampling frequency. A compare and combine circuit may be provided to combine the simulated response with the measured response using algorithms such as, without limitation, weighted average and theoretical control loop that drives the simulated response towards the measured response. Simulation of the physical system takes in account of properties like mass, inertial, viscous friction, inductance resistance, etc. to predict what the states and outputs of the physical system will be by knowing the input. In one aspect, the controller 7004 may be a single core or multicore controller LM4F230H5QR as described in connection with FIGS. 15-17B.

[0335] In one aspect, the driver 7010 may be a A3941 available from Allegro Microsystems, Inc. The A3941 driver 7010 is a full-bridge controller for use with external N-channel power metal oxide semiconductor field effect transistors (MOSFETs) specifically designed for inductive loads, such as brush DC motors. The driver 7010 comprises a unique charge pump regulator provides full (>10 V) gate drive for battery voltages down to 7 V and allows the A3941 to operate with a reduced gate drive, down to 5.5 V. A bootstrap capacitor may be employed to provide the above-battery supply voltage required for

N-channel MOSFETs. An internal charge pump for the high-side drive allows DC (100% duty cycle) operation. The full bridge can be driven in fast or slow decay modes using diode or synchronous rectification. In the slow decay mode, current recirculation can be through the high-

⁵ cay mode, current recirculation can be through the highside or the lowside FETs. The power FETs are protected from shoot-through by resistor adjustable dead time. Integrated diagnostics provide indication of undervoltage, overtemperature, and power bridge faults, and can be

¹⁰ configured to protect the power MOSFETs under most short circuit conditions. Other motor drivers may be readily substituted for use in the absolute positioning system 7000. Accordingly, the present disclosure should not be limited in this context.

¹⁵ [0336] Having described a general architecture for implementing various aspects of an absolute positioning system 7000 for a sensor arrangement 7002, the disclosure now turns to FIGS. 101-103 for a description of one aspect of a sensor arrangement for the absolute positioning system 7000. In the aspect illustrated in FIG. 101,

the sensor arrangement 7002 comprises a position sensor 7100, a magnet 7102 sensor element, a magnet holder 7104 that turns once every full stroke of the longitudinally-movable drive member 1111 (FIG. 100), and a gear

assembly 7106 to provide a gear reduction. A structural element such as bracket 7116 is provided to support the gear assembly 7106, the magnet holder 7104, and the magnet 7102. The position sensor 7100 comprises one or more than one magnetic sensing elements such as
Hall elements and is placed in proximity to the magnet 7102. Accordingly, as the magnet 7102 rotates, the magnetic sensing elements of the position sensor 7100 determine the absolute angular position of the magnet 7102 over one revolution.

³⁵ [0337] In various aspects, any number of magnetic sensing elements may be employed on the absolute positioning system 7000, such as , for example, magnetic sensors classified according to whether they measure the total magnetic field or the vector components of the

40 magnetic field. The techniques used to produce both types of magnetic sensors encompass many aspects of physics and electronics. The technologies used for magnetic field sensing include search coil, fluxgate, optically pumped, nuclear precession, SQUID, Hall-effect, aniso-

⁴⁵ tropic magnetoresistance, giant magnetoresistance, magnetic tunnel junctions, giant magnetoimpedance, magnetostrictive/piezoelectric composites, magnetodiode, magnetotransistor, fiber optic, magnetooptic, and microelectromechanical systems-based magnetic sensors, among others.

[0338] In the illustrated aspect, the gear assembly 7106 comprises a first gear 7108 and a second gear 7110 in meshing engagement to provide a 3:1 gear ratio connection. A third gear 7112 rotates about shaft 7114. The third gear is in meshing engagement with the longitudinally-movable drive member 1111 and rotates in a first direction as the longitudinally-movable drive member 1111 advances in a distal direction D and rotates in a

second direction as the longitudinally-movable drive member 1111 retracts in a proximal direction P. The second gear 7110 also rotates about the shaft 7114 and therefore, rotation of the second gear 7110 about the shaft 7114 corresponds to the longitudinal translation of the longitudinally-movable drive member 1111. Thus, one full stroke of the longitudinally-movable drive member 1111 in either the distal or proximal directions D, P corresponds to three rotations of the second gear 7110 and a single rotation of the first gear 7108. Since the magnet holder 7104 is coupled to the first gear 7108, the magnet holder 7104 makes one full rotation with each full stroke of the longitudinally-movable drive member 1111.

[0339] FIG. 102 is an exploded perspective view of the sensor arrangement 7002 for the absolute positioning system 7000 showing a circuit 1106 and the relative alignment of the elements of the sensor arrangement 7002, according to one aspect. The position sensor 7100 (not shown in this view) is supported by a position sensor holder 7118 defining an aperture 7120 suitable to contain the position sensor 7100 in precise alignment with a magnet 7102 rotating below. The fixture is coupled to the bracket 7116 and to the circuit 1106 and remains stationary while the magnet 7102 rotates with the magnet holder 7104. A hub 7122 is provided to mate with the first gear 7108 and the magnet holder 7104.

[0340] FIG. 103 is a schematic diagram of one aspect of a position sensor 7100 sensor for an absolute positioning system 7000 comprising a magnetic rotary absolute positioning system, according to one aspect. In one aspect, the position sensor 7100 may be implemented as an AS5055EQFT single-chip magnetic rotary position sensor available from Austria Microsystems, AG. The position sensor 7100 is interfaced with the controller 7004 to provide an absolute positioning system 7000. The position sensor 7100 is a low voltage and low power component and includes four Hall-effect elements 7128A, 7128B, 7128C, 7128D in an area 7130 of the position sensor 7100 that is located above the magnet 7102 (FIGS. 99, 100). A high resolution ADC 7132 and a smart power management controller 7138 are also provided on the chip. A CORDIC processor 7136 (for COordinate Rotation DIgital Computer), also known as the digit-by-digit method and Volder's algorithm, is provided to implement a simple and efficient algorithm to calculate hyperbolic and trigonometric functions that require only addition, subtraction, bitshift, and table lookup operations. The angle position, alarm bits and magnetic field information are transmitted over a standard serial communication interface such as an SPI interface 7134 to the controller 7004. The position sensor 7100 provides 12 or 14 bits of resolution. The position sensor 7100 may be an AS5055 chip provided in a small QFN 16-pin 4x4x0.85mm package.

[0341] The Hall-effect elements 7128A, 7128B, 7128C, 7128D are located directly above the rotating magnet. The Hall-effect is a well known effect and will

not be described in detail herein for the sake of conciseness and clarity of disclosure. Generally, the Hall-effect is the production of a voltage difference (the Hall voltage) across an electrical conductor, transverse to an electric current in the conductor and a magnetic field perpendicular to the current. It was discovered by Edwin Hall in 1879. The Hall coefficient is defined as the ratio of the induced electric field to the product of the current density

and the applied magnetic field. It is a characteristic of the
 material from which the conductor is made, since its value
 depends on the type, number, and properties of the
 charge carriers that constitute the current. In the AS5055
 position sensor 7100, the Hall-effect elements 7128A,
 7128B, 7128C, 7128D are capable producing a voltage

¹⁵ signal that is indicative of the absolute position of the magnet 7102 (FIGS. 186, 187) in terms of the angle over a single revolution of the magnet 7102. This value of the angle, which is unique position signal, is calculated by the CORDIC processor 7136 is stored onboard the

20 AS5055 position sensor 7100 in a register or memory. The value of the angle that is indicative of the position of the magnet 7102 over one revolution is provided to the controller 7004 in a variety of techniques, e.g., upon power up or upon request by the controller 7004.

²⁵ [0342] The AS5055 position sensor 7100 requires only a few external components to operate when connected to the controller 7004. Six wires are needed for a simple application using a single power supply: two wires for power and four wires 7140 for the SPI interface 7134 with

30 the controller 7004. A seventh connection can be added in order to send an interrupt to the controller 7004 to inform that a new valid angle can be read.

[0343] Upon power-up, the AS5055 position sensor 7100 performs a full power-up sequence including one angle measurement. The completion of this cycle is indicated as an INT output 7142 and the angle value is stored in an internal register. Once this output is set, the AS5055 position sensor 7100 suspends to sleep mode. The controller 7004 can respond to the INT request at the INT output 7142 by reading the angle value from the AS5055 position sensor 7100 over the SPI interface

7134. Once the angle value is read by the controller 7004, the INT output 7142 is cleared again. Sending a "read angle" command by the SPI interface 7134 by the con⁴⁵ troller 7004 to the position sensor 7100 also automatically

powers up the chip and starts another angle measurement. As soon as the controller 7004 has completed reading of the angle value, the INT output 7142 is cleared and a new result is stored in the angle register. The completion of the angle measurement is again indicated by setting the INT output 7142 and a corresponding flag in the

status register. **[0344]** Due to the measurement principle of the AS5055 position sensor 7100, only a single angle measurement is performed in very short time (\sim 600µs) after each power-up sequence. As soon as the measurement of one angle is completed, the AS5055 position sensor 7100 suspends to power-down state. An on-chip filtering

of the angle value by digital averaging is not implemented, as this would require more than one angle measurement and consequently, a longer power-up time which is not desired in low power applications. The angle jitter can be reduced by averaging of several angle samples in the controller 7004. For example, an averaging of 4 samples reduces the jitter by 6dB (50%).

[0345] As discussed above, the electric motor 1102 positioned within the handle 1042 of surgical instrument system 1000 can be utilized to advance and/or retract the firing system of the shaft assembly 1200, including firing members 1272 and 1280, for example, relative to the end effector 1300 of the shaft assembly 1200 in order to staple and/or incise tissue captured within the end effector 1300. In various circumstances, it may be desirable to advance the firing members 1272 and 1280 at a desired speed, or within a range of desired speeds. Likewise, it may be desirable to retract the firing members 1272 and 1280 at a desired speed, or within a range of desired speeds. In various circumstances, the controller 7004 of the handle 1042, for example, and/or any other suitable controller, can be configured to control the speed of the firing members 1272 and 1280. In some circumstances, the controller can be configured to predict the speed of the firing members 1272 and 1280 based on various parameters of the power supplied to the electric motor 1102, such as voltage and/or current, for example, and/or other operating parameters of the electric motor 1102. The controller can also be configured to predict the current speed of the firing members 1272 and 1280 based on the previous values of the current and/or voltage supplied to the electric motor 1102, and/or previous states of the system like velocity, acceleration, and/or position. Furthermore, the controller can also be configured to sense the speed of the firing members 1272 and 1280 utilizing the absolute positioning sensor system described above, for example. In various circumstances, the controller can be configured to compare the predicted speed of the firing members 1272 and 1280 and the sensed speed of the firing members 1272 and 1280 to determine whether the power to the electric motor 1102 should be increased in order to increase the speed of the firing members 1272 and 1280 and/or decreased in order to decrease the speed of the firing members 1272 and 1280. U.S. Patent No. 8,210,411, entitled MOTOR-DRIVEN SURGICAL CUTTING INSTRUMENT, which is incorporated herein by reference in its entirety. U.S. Patent No. 7,845,537, entitled SURGICAL INSTRUMENT HAVING RECORDING CAPABILITIES, which is incorporated herein by reference in its entirety.

[0346] Using the physical properties of the instruments disclosed herein, turning now to FIGS. 104 and 105, a controller, such as controller 7004, for example, can be designed to simulate the response of the actual system of the instrument in the software of the controller. The simulated response is compared to a (noisy and discrete) measured response of the actual system to obtain an "observed" response, which is used for actual feedback

decisions. The observed response is a favorable, tuned, value that balances the smooth, continuous nature of the simulated response with the measured response, which can detect outside influences on the system. With regard to FIGS. 104 and 105, a firing element, or cutting element, in the end effector 1300 of the shaft assembly 1200 can be moved at or near a target velocity, or speed. The systems disclosed in FIGS. 102 and 103 can be utilized to

move the cutting element at a target velocity. The systems can include a feedback controller 4200, which can be one of any feedback controllers, including, but not limited to a PID, a State Feedback, LQR, and/or an Adaptive controller, for example. The systems can further include a power source. The power source can convert the

¹⁵ signal from the feedback controller 4200 into a physical input to the system, in this case voltage, for example. Other examples include, but are not limited to, pulse width modulated (PWM) voltage, frequency modulated voltage, current, torque, and/or force, for example.

20 [0347] With continued reference to FIGS. 104 and 105, the physical system referred to therein is the actual drive system of the instrument configured to drive the firing member, or cutting member. One example is a brushed DC motor with gearbox and mechanical links to an artic-

²⁵ ulation and/or knife system. Another example is the electric motor 1102 disclosed herein that operates the firing member 10060 and the articulation driver 10030, for example, of an interchangeable shaft assembly. The outside influence 4201 referred to in FIGS. 104 and 105 is
³⁰ the unmeasured, unpredictable influence of things like tissue, surrounding bodies and friction on the physical system, for example. Such outside influence can be re-

ferred to as drag and can be represented by a motor 4202 which acts in opposition to the electric motor 1102, for
example. In various circumstances, outside influence, such as drag, is the primary cause for deviation of the simulation of the physical system from the actual physical system. The systems depicted in FIGS. 104 and 105 and further discussed below can address the differences be-

40 tween the predicted behavior of the firing member, or cutting member, and the actual behavior of the firing member, or cutting member.

[0348] With continued reference to FIGS. 104 and 105, the discrete sensor referred to therein measures physical 45 parameters of the actual physical system. One aspect of such a discrete sensor can include an absolute positioning sensor and system described herein, such as the magnet 7102. As the output of such a discrete sensor can be a digital signal (or connected to a digital data 50 acquisition system) its output may have finite resolution and sampling frequency. The output of the discrete sensor can be supplied to a controller, such as controller 7004, for example. In various circumstances, the controller can combine the simulated, or estimated, response 55 with the measured response. In certain circumstances, it may be useful to use enough measured response to ensure that the outside influence is accounted for without making the observed response unusably noisy. Exam-

ples for algorithms that do so include a weighted average and/or a theoretical control loop that drives the simulated response towards the measured response, for example. Ultimately, further to the above, the simulation of the physical system takes in account of properties like mass, inertial, viscous friction, and/or inductance resistance, for example, to predict what the states and outputs of the physical system will be by knowing the input. FIG. 103 shows an addition of evaluating and measuring the current supplied to operate the actual system, which is yet another parameter that can be evaluated for controlling the speed of the cutting member, or firing member, of the shaft assembly 1200, for example. By measuring current in addition to or in lieu of measuring the voltage, in certain circumstances, the physical system can be made more accurate. Nonetheless, the ideas disclosed herein can be extended to the measurement of other state parameters of other physical systems.

[0349] FIG. 106 shows a handle assembly 6302 of a modular surgical instrument, similar to surgical instrument 10 shown and described with regard to FIGS. 1-105 and more particularly in regard to FIGS. 1-14. The handle assembly 6302 may be engaged with an interchangeable shaft assembly (not shown), such as shaft assembly 200 described herein in FIGS. 1-14. The handle assembly 6302 includes a closure trigger 6304, pistol grip portion 6306, an interface 6308, and a user display 6310, which may also be configured to receive an input from a user. The handle assembly 6302 also includes a closure release button assembly 6212, which can be depressed by the operator to release a closure lock thereby returning an end effector (not shown) to a desired configuration. The handle assembly 6302 is shown with a socket 6314 for a battery and the battery pack is removed.

[0350] The interface 6308 may comprise at least one user-actuated input device, such as a switch, that may be configured to provide functionality as described herein. According to aspects the interface 6308 may be utilized to implement an articulation function of an interchangeable shaft assembly. Additionally, according to the aspect shown in FIG. 106, the interface 6308 comprises a plurality of user-actuated input devices, switches 6316, 6318, 6320, which can be utilized by the operator, in part, to activate and control functions associated with an end effector of an interchangeable shaft assembly, as described herein. According to an aspect, the handle assembly 6302 may be combined with an interchangeable shaft assembly having a cutting member as described herein. In such an aspect, the switches 6316, 6318, 6320 may be used to control activation and speed of the cutting member.

[0351] As shown in FIG. 106, the switch 6316 is a button style switch and switches 6318 and 6320 are part of a toggle style switch. The interface 108 may be configured such that the switch 6316 is dedicated to activation and/or selection of the speed of a cutting member. Further, switch 6318 may be dedicated to increasing the speed of the cutting member and the switch 6320 may

be dedicated to decreasing the speed of the cutting member. Other means for controlling the switches 6318, 6320 and/or replacing functionality of the switches 6318, 6320 with a single switch device are also contemplated within the scope of the present disclosure.

[0352] Furthermore, FIG. 107 illustrates a schematic block diagram of a control system 6400 for a surgical instrument that comprises the handle assembly 6302 of FIG. 106. The interface 6308 of the handle assembly

¹⁰ 6302 may be used to operate a cutting member of an interchangeable shaft assembly engaged with handle assembly 6302 via the control system 6400 of FIG. 107. The control system 6400 comprises a controller 6402 which can be configured to receive an input signal via

¹⁵ interface 6405, which corresponds to the interface 6308 shown in FIG. 106, and, in response, activates the motor 6404 to cause a cutting member of an end effector to move in accordance with such an input signal. Examples of suitable controllers are described elsewhere in this

²⁰ document and in particular in connection with FIGS. 15-17B, among others. Each of the switches 6406A, 6406B, 6406C may be coupled to or correspond to contacts of each of the switches 6316, 6318, 6320, respectively, shown in FIG. 106. The switches 6406A-C are cou-

²⁵ pled to the controller 6402 via one of three electrical circuits 6408A-C, respectively. The reader will appreciate that other combinations of switches and circuits can be utilized in conjunction with the interface 108.

[0353] Further, the controller 6402 may comprise a 30 processor 6410 and/or one or more memory units 6412. By executing instruction code stored in the memory 6412, the processor 6410 may control and communicate with various components of the surgical instrument, such as the motor 6404 and/or a user display 6414. The controller

³⁵ 6408 may be implemented using integrated and/or discrete hardware elements, software elements, and/or a combination of both. Examples of integrated hardware elements may include processors, microprocessors, microcontrollers, integrated circuits, application specific integrated circuits (ASIC), programmable logic devices

(PLD), digital signal processors (DSP), field programmable gate arrays (FPGA), logic gates, registers, semiconductor devices, chips, microchips, chip sets, microcontroller, system-on-chip (SoC), and/or system-in-package

⁴⁵ (SIP). Examples of discrete hardware elements may include circuits and/or circuit elements (e.g., logic gates, field effect transistors, bipolar transistors, resistors, capacitors, inductors, relay and so forth). In other aspects, the controller 6402 may include a hybrid circuit compris⁵⁰ ing discrete and integrated circuit elements or components on one or more substrates, for example. The controller 6402 may comprise an internal analog-to-digital (A/D) converter or maybe coupled to an external A/D converter 6424 to digitize analog signals from various sourc⁵⁵ es, such as, for example, variable resistor 6423, generally referred to as a potentiometer.

[0354] The memory 6414 may be used to store instructions that represent an operating mode of the surgical

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instrument, such as for example, an algorithm for implementing an operating mode of the electric motor 6404 of the surgical instrument. The operating mode may define an operation to be performed by the surgical instrument. The operating mode may pertain to the operation of a cutting member of an end effector that is controlled by the electric motor 6404. In one aspect, the operating mode may be associated with a parameter sensed by sensor 6416 and/or sensor 6418. In one aspect, the controller 6402 may be configured to implement a pre-determined operation of the electric motor 6416 when at least one parameter sensed by the sensor 6416 and/or sensor 6418 meets or exceeds a predefined threshold value according to the operating mode for the electric motor 6416. One or both of the sensors 6416, 6418 may be configured to sense a different parameter or the same parameter associated with an aspect and/or a component of the surgical instrument. The sensors 6416, 6418 may sense the same parameter at different locations of the surgical instrument. According to aspects, one or both of the sensors 6416, 6418 may be configured to sense a parameter associated with the electric motor, a drive system, a component of an end effector, or combinations thereof. Furthermore, the operating mode may be implemented according to a predetermined threshold of the parameter sensed by either or both sensors 6416, 6418. In one aspect, the sensor 6416 may sense current supplied by the electric motor, and may decrease power to the electric motor 6404 when a predetermined current threshold has been reached. In other aspects, other parameters may sensed by one or both of the sensors 6416, 6418, for example, parameters including tissue parameters of a tissue section clamped within an end effector as described herein.

[0355] Furthermore, the motor controller 6420 in operable communication with the controller 6402. The motor controller 6420 can be configured to control a direction and/or speed of rotation of the motor 6404. For example, the motor 6404 can be powered by a battery such as, for example, the battery 6422 and the motor controller 6420 may be configured to determine the voltage polarity applied to the motor 6404 by the battery 6422 and, in turn, the direction of rotation of the motor 6404 based on input from the controller 6402. For example, the motor 6404 may reverse the direction of its rotation from a clockwise direction to a counterclockwise direction when the voltage polarity applied to the motor 6404 by the battery 6422 is reversed by the motor controller 6402 based on an input from the controller 6402. Examples of suitable motor controllers are described elsewhere in this document. [0356] With reference now to FIGS. 108-111 a detail view of one aspect of the interface 6308 is shown. The switches 6316, 6318, 6320 correspond to the switches 6406A, 6406B, 6406C shown in FIG. 107 and can each comprise open-biased dome switches as illustrated. Other types of switches can also be employed such as, for example, capacitive switches. In other aspects, the switches 6406A, 6406B, 6406C may all be of the same type or different types. For example, in one aspect switch 6406A may be a capacitive switch and switches 6406B, 6406C may be open-biased dome switches. As illustrated in FIGS. 110, 111, the dome switches 6318, 6320 are controlled by a toggle or rocker 6322 and dome the switch 6316 is controlled by a button 6324. Other mechanisms for controlling the switches 6318, 6320 and/or replacing functionality of switches 6318, 6320 with a single switch device are also contemplated within the scope of the present disclosure.

[0357] In the neutral position, illustrated in FIG. 110, both of the switches 6308, 6320 are biased in the open position. The operator, for example, may increase the speed of a cutting member coupled to the motor 6404

¹⁵ (FIG. 107) by tilting the rocker forward thereby depressing the dome switch 6318, as illustrated in FIG. 111. In result, the circuit 6408B of FIG. 107 may be closed signaling the controller 6402 to activate the motor 6404 to increase the speed of the cutting member, as described

²⁰ above. The motor 6404 may continue to increase the speed of the cutting member until the operator releases the rocker 6322 thereby allowing the switch 6406B to return to the open position and the rocker 6322 to the neutral position (as shown in FIG. 110). In some circum-

stances, the controller 6402 may be able to identify when the cutting member has reached a predetermined maximum speed and, at such point, apply constant power to the motor 6404 regardless of whether the switch 6406B is being depressed. Alternatively, the operator may de-

³⁰ crease the speed of the cutting member by tilting the rocker 6322 back thereby depressing the switch 6406C, for example. In result, the circuit 6408C may be closed signaling the controller 6402 to reduce power to the motor 6404 to reduce the speed of the cutting member, as de-

³⁵ scribed above. The motor 6404 may continue to decrease the speed of the cutting member of the end effector until the operator releases the rocker 6322 thereby allowing the switch 6406C to return to the open position and the rocker 6322 to the neutral position (as shown in

40 FIG. 111). In some circumstances, the controller 6402 may be able to identify when the end effector has reached a predetermined minimum speed, at such point, apply constant power to the motor 6404 regardless of whether the switch 6406C is being depressed.

⁴⁵ [0358] In aspects, the control system 6400 may include a virtual detent that may alert the operator when the cutting member of the end effector reaches a particular maximum or minimum speed. For example, the operator may tilt the rocker 6322 to increase or decrease the speed of

⁵⁰ a cutting member of an end effector. Upon reaching the maximum or minimum speed, the controller 6402 may stop increasing or decreasing the speed of the cutting member of the end effector. Alternatively, a mechanical detent can also be used to provide haptic feedback for ⁵⁵ the operator that the cutting member reached a maximum or minimum speed. Other forms of feedback may be utilized such as audio feedback, for example.

[0359] According to aspects and with reference to FIG.

106, the switches 6318, 6320 of handle assembly 6302 may be replaced by a single user-actuated input device. As shown in FIG. 112, the interface 6508 of the handle assembly 6502 comprises a rotary potentiometer 6504 that may be used to adjust the speed of cutting member. For example, in one aspect, counter-clockwise rotation of the rotary potentiometer 6504 increases the speed of the cutting member whereas clockwise rotation of the rotary potentiometer 6504 decreases the speed of the cutting member. The output of the rotary potentiometer 6504 can be provided to the external A/D converter 6424 or to the internal A/D converter of the controller 6402 of FIG. 107. In other aspects, the rotary potentiometer 6504 may be replaced with a rotary encoder whose output can be provided to the controller 6402.

[0360] In addition, in the aspect shown in FIG. 113, the interface 6608 of the handle assembly 6602 comprises a slide potentiometer 6604 that may be used to adjust the speed of cutting member such that sliding the sliding bar to the left increases the speed of the cutting member whereas sliding the sliding bar to the right decreases the speed of the cutting member. The output of the slide potentiometer 6604 can be provided to the external A/D converter 6424 or to the internal A/D converter of the controller 6402 of FIG. 107. In other aspects, the slide potentiometer 6604 may be replaced with a slide encoder whose output can be provided to the controller 6402.

[0361] According to aspects and with reference to FIG. 107 and FIGS. 112, 113, the rotary potentiometer 6504 or the slide potentiometer 6604 may be used to replace the functionality associated with switches 6406B, 6406C. Further, referring to FIG. 107 and FIGS. 112, 113, the circuits 6408B, 6408C may be combined into a single circuit that is associated with rotary potentiometer 6504 or slide potentiometer 504. As shown in FIG. 107, in one aspect, the rotary potentiometer 6504 or slide potentiometer 6604 may comprise a variable resistor 6423 that is coupled to the A/D converter 6424. Accordingly, the controller 6402 may receive a digitized signal from either the rotary potentiometer 6504 or slide potentiometer 6604 and provide it to the motor controller 6420 for controlling the speed of motor 6404.

[0362] According to one aspect, the switches 6316, 6318, 6320 of handle assembly 6302 may be configured so that by pressing one of the switches, an operating mode of the surgical instrument 10, of which the handle assembly 6302 is a component, may be overridden. Referring back to FIG. 107, the controller 6402 may be configured to receive an electrical signal that corresponds to one of switches 6406A, 6406B, 6406C of interface 6405 being activated. Upon receiving the electrical signal, the controller 6402 may override an operating mode of the surgical instrument such that a user is then able to control aspects of the surgical instrument using interface 205. In one aspect, switch 6406A is a button that acts as an activation switch that overrides an operating mode associated with a component of an end effector, such as a cutting member or an articulation member.

Accordingly, the controller 6402 may implement an operating mode of an electric motor of the surgical instrument stored in memory 6412, where the operating mode comprises the motor 6404 moving the cutting member

⁵ or the articulation member. Once switch 6406A is activated, the operating mode of the electric motor 6404 that moves a component of the surgical instrument, such as, for example, a cutting member or an articulation member is overridden.

10 [0363] Once the controller has overridden the operating mode, the controller 6402 may then be able to receive inputs from switches 6406B, 6406C and operate the component of the surgical instrument 10, such as, for example, a cutting member or an articulation member via the

¹⁵ motor 6404. According to one aspect, an input based on switch 206B may increase the speed of the cutting member and an input based on the switch 6406C may decrease the speed of the cutting member. In one aspect, once the controller has overridden the operating mode,

20 the controller may be able to receive inputs from the switches 6406B, 6406C and operate the cutting member via the motor 6404 when at least one parameter sensed by one or both of the sensors 6416, 6418 meets or exceeds a predefined threshold value.

²⁵ [0364] The operating mode of the motorized surgical instrument described herein may be based on parameters sensed by one or both of sensors 6416, 6418. In one aspect, the operating mode may be based on tissue parameters of tissue engaged by an end effector, of which

the cutting member is a component. In addition, the operating mode may be further or instead based on a parameter associated with a drive system or other component, such as a cutting member or an articulation member, of the surgical instrument as described herein. Fur-

thermore, in other aspects, additional sensors to sensors 6416, 6418 may be added and those sensors may sense any parameters as described herein. In one aspect, the controller 6402 may be configured to implement the operating mode so that is adjusts power to the electric motor

40 6404 if a parameter sensed by one or both of sensors 6416, 6418 exceeds a predefined threshold value for that parameter.

[0365] According to one aspect, the switch 6406A may be activation switch that starts the override process. In

another aspect, more than one switch 6406A, 6406B, 6406C may be activated to start the override process. In one aspect, a secondary button may provided that is activated in combination with regular control switch, such as one of switch 6406A, 6406B, 6406C, such that activation of the secondary button in combination with the regular control switch drives the control system 200 to stop automatic modification of motions related to the regular control switch.

[0366] In aspects, the user display 6310, and correspondingly user display 6414 shown in FIG. 107, may comprise an input device electrically coupled to the controller 6402 such that a function of the surgical instrument may be adjusted by a user input prior to use of the surgical instrument. In one aspect, the user display 6310 of handle assembly 6302 may comprise an input device that allows a user to select an operating mode or other functionality of the surgical instrument as a form of setup control. The operating mode may be selected from a plurality of operating modes stored in memory 6412. Controller 6402 may be configured to receive an input from the user display 6414, prior to beginning an operating mode, where the input comprises a command that configures the controller 6402 to override the operating mode for the electric motor 6404 upon receiving the activation of switch 6406A. In other aspects, the controller may be configured to override the operating mode for the electric motor upon receiving the activation of any one or a combination of switches 6406A, 6406B, 6406C.

[0367] As shown in FIG. 114 and with reference also to FIG. 107 a method 6700 for operating a surgical instrument 10, similar to surgical instrument 10 shown and described with regard to FIGS. 1-105 and more particularly in regard to FIGS. 1-14, comprising a handle assembly 6302, 6502, 6602. The method 6700 comprises sensing 6702 a parameter, via a sensor 6416, 6418 of or associated with the control system 6400 of the surgical instrument 10, implementing 6704, via a controller 6402 of the surgical instrument 10, an operating mode of an electric motor 6404 of the surgical instrument 10, receiving 6706, via the controller 6402, an electrical signal, and overriding 6708, via the controller 6402, the operating mode of the electric motor 6404 upon receiving 6706 the electrical signal. The parameter may be associated with a drive system of the surgical instrument 10 and the electrical signal comprises an activation of at least one useractuated input device electrically coupled to the controller 6402. In addition, the operating mode of the surgical instrument 10 may be based on the parameter sensed by the sensor.

[0368] Further, in one aspect, implementing the operating mode of the electric motor 6404 comprises the controller 6402 adjusting power to the electric motor 6404 if the parameter exceeds a predefined threshold value. In another aspect, the method 6700 further comprises receiving, via an input from a user selection display of the surgical instrument, a command that configures the controller 6402 to override the operating mode for the electric motor 6404 upon receiving the activation of the at least one user-actuated input device. In yet another aspect, the method 6700 further comprises operating the electric motor 6404 according to an input received via the at least one user-actuated input device upon the controller 6402 overriding the operating system of electric motor 6404. [0369] With reference to FIG. 106 and FIGS. 1-14, the handle assembly 6302 may be also used with an interchangeable shaft assembly 200 that has an end effector 300 with a cutting member (knife bar 280) along with an articulation member (articulation driver 230) to allow for multiple functionalities for the end effector 300. It may be desirable to prevent not only two contemporaneous signals for activating the two functionalities of the end effector at the same time, i.e., the cutting member and the articulation member, it may also be desirable to allow a first functionality of the end effector 300, such as the cutting member, to complete an operating mode associated

⁵ with the first functionality prior to activating a second functionality of the end effector, such as an articulation member. Furthermore, in the event that completion of the operating mode associated with the first functionality may be expedited, it may be desirable to do so.

10 [0370] With reference to FIG. 107, in one aspect, the controller 6402 may be configured to implement an operating mode of the surgical instrument 10, where the operating mode comprises operating a first functionality of a component of the surgical instrument 10, such as a

¹⁵ cutting member or articulation member, according to a first speed. The controller 6402 may be configured to receive an electrical signal associated with an activation of at least one user-actuated input device, such as one or more of switches 6406A-C, and the controller 6402

²⁰ may be configured to override the operating mode of the surgical instrument 10 and to operate the first functionality of the surgical instrument, such as the cutting member, according to a second speed that is greater than the first speed. In one aspect, the electric motor 6404 may

25 be operated to cause a cutting member to retract from an extended position to a home or base position at a speed that is greater than a predetermined speed provided for in the operating mode at which the cutting member is supposed to retract. In another aspect, the electric 30 motor 6404 may be operated to cause the cutting member to extend distally at a speed that is greater than a predetermined speed provided for in the operating mode at which the cutting member is supposed to extend. In other aspects, the controller 6402 may be configured to 35 wait until an operation of the functionality of an end effector, for example the cutting member, is complete, prior to changing the speed of the functionality. Accordingly, in one aspect, the controller 6402 may be configured to wait until a transection operation is complete prior to ad-

vancing the speed of a cutting member. [0371] In addition, the controller 6402 may be configured to implement a second operating mode for a second functionality of a component of the surgical instrument 10 once the operation of a first operating mode is com-

⁴⁵ plete. In one aspect, once the electric motor 6404 causes a cutting member to return to a base position following the completion of an operating mode associated with the cutting member, for example extension of the cutting member during transection, the controller may then im-

⁵⁰ plement an operating mode of the articulation member. Similarly to the above, the surgical instrument 10 may include at least one sensor configured to sense a parameter associated with the surgical instrument 10 and the operating mode of the functionality of the surgical instru-⁵⁵ ment 10 is based on the parameter sensed by the at least one sensor 6416, 6418. Accordingly, the speed of a component of the surgical instrument 10, such as a cutting member or an articulation member, may be adjusted

based on the parameter sensed by the at least one sensor 6416, 6418.

[0372] As shown in FIG. 115 and with reference also to FIG. 107 a method 6800 for operating a surgical instrument 10, similar to surgical instrument 10 shown and described with regard to FIGS. 1-105 and more particularly in regard to FIGS. 1-14, comprising a handle assembly 6302, 6502, 6602. The method 6800 comprises implementing 6802, via a control system 6400 of the surgical instrument 10 comprising controller 6402, an operating mode of the surgical instrument 10, where the operating mode comprises operating a component of the surgical instrument 10 according to a first speed, receiving 6804, via the controller 6402, an electrical signal associated with an activation of at least one user-actuated input device electrically coupled to the controller 6402, overriding 6806, via the controller 6402, the operating mode of the surgical instrument 10 upon receiving the electrical signal, and operating 6808, via the controller 6402, the component of the surgical instrument 10 according to a second speed that is greater than the first speed. According to various aspects of the present disclosure, the surgical instrument 10 may have a first operating mode and a second operating mode each associated with a first component and a second component, respectively such that the method further comprises implementing, via the controller 6402 of the surgical instrument 10, a second operating mode of the surgical instrument 10. In one aspect of the present disclosure, the first component may be a cutting member (knife bar 280) and the second component may be an articulation member (articulation driver 230). In another aspect of the present disclosure, the method further comprises sensing a parameter, via at least one sensor 6416, 6418 of the surgical instrument 10. The parameter may be associated with a drive system of the surgical instrument 10. In addition, either the first operating mode and/or the second operating made may be based on the parameter sensed by the at least one sensor 6416, 6418.

[0373] In another aspect of the present disclosure, operating 6808 the component of the surgical instrument 10 according to the second speed comprises operating the component to return the component, for example a cutting member or an articulation member, of the surgical instrument 10 to a base position. Furthermore, in one aspect, overriding 6806 the operating mode comprises completing an operation associated the component, for example a cutting member or an articulation member, of the surgical instrument 10 prior to operating the component of the surgical instrument 10 prior to example a cutting the surgical instrument 10 prior to preating the component of the surgical instrument 10 according to the second speed.

[0374] Although the various aspects of the devices have been described herein in connection with certain disclosed aspects, many modifications and variations to those aspects may be implemented. For example, different types of end effectors may be employed. Also, where materials are disclosed for certain components, other materials may be used. The foregoing description and

following claims are intended to cover all such modification and variations.

[0375] Any patent, publication, or other disclosure material, in whole or in part, that is said to be incorporated
⁵ by reference herein is incorporated herein only to the extent that the incorporated materials does not conflict with existing definitions, statements, or other disclosure material set forth in this disclosure. As such, and to the extent necessary, the disclosure as explicitly set forth

¹⁰ herein supersedes any conflicting material incorporated herein by reference. Any material, or portion thereof, that is said to be incorporated by reference herein, but which conflicts with existing definitions, statements, or other disclosure material set forth herein will only be incorpo-

¹⁵ rated to the extent that no conflict arises between that incorporated material and the existing disclosure material.

[0376] While various details have been set forth in the foregoing description, it will be appreciated that the various aspects of the motorized surgical instruments may be practiced without these specific details. For example, for conciseness and clarity selected aspects have been shown in block diagram form rather than in detail. Some portions of the detailed descriptions provided herein may

²⁵ be presented in terms of instructions that operate on data that is stored in a computer memory. Such descriptions and representations are used by those skilled in the art to describe and convey the substance of their work to others skilled in the art. In general, an algorithm refers ³⁰ to a self-consistent sequence of steps leading to a de-

sired result, where a "step" refers to a manipulation of physical quantities which may, though need not necessarily, take the form of electrical or magnetic signals capable of being stored, transferred, combined, compared,

 and otherwise manipulated. It is common usage to refer to these signals as bits, values, elements, symbols, characters, terms, numbers, or the like. These and similar terms may be associated with the appropriate physical quantities and are merely convenient labels applied to
 these quantities.

[0377] Although various aspects have been described herein, many modifications, variations, substitutions, changes, and equivalents to those aspects may be implemented and will occur to those skilled in the art. Also,

⁴⁵ where materials are disclosed for certain components, other materials may be used. It is therefore to be understood that the foregoing description and the appended claims are intended to cover all such modifications and variations as falling within the scope of the disclosed aspects. The following claims are intended to cover all such

modification and variations. **[0378]** In a general sense, those skilled in the art will recognize that the various aspects described herein which can be implemented, individually and/or collectively, by a wide range of hardware, software, firmware, or any combination thereof can be viewed as being composed of various types of "electrical circuitry." Conse-

quently, as used herein "electrical circuitry" includes, but

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is not limited to, electrical circuitry having at least one discrete electrical circuit, electrical circuitry having at least one integrated circuit, electrical circuitry having at least one application specific integrated circuit, electrical circuitry forming a general purpose computing device configured by a computer program (e.g., a general purpose computer configured by a computer program which at least partially carries out processes and/or devices described herein, or a processor configured by a computer program which at least partially carries out processes and/or devices described herein), electrical circuitry forming a memory device (e.g., forms of random access memory), and/or electrical circuitry forming a communications device (e.g., a modem, communications switch, or optical-electrical equipment). Those having skill in the art will recognize that the subject matter described herein may be implemented in an analog or digital fashion or some combination thereof.

[0379] The foregoing detailed description has set forth various aspects of the devices and/or processes via the use of block diagrams, flowcharts, and/or examples. Insofar as such block diagrams, flowcharts, and/or examples contain one or more functions and/or operations, it will be understood by those within the art that each function and/or operation within such block diagrams, flowcharts, or examples can be implemented, individually and/or collectively, by a wide range of hardware, software, firmware, or virtually any combination thereof. In one aspect, several portions of the subject matter described herein may be implemented via Application Specific Integrated Circuits (ASICs), Field Programmable Gate Arrays (FPGAs), digital signal processors (DSPs), or other integrated formats. Those skilled in the art will recognize, however, that some aspects of the aspects disclosed herein, in whole or in part, can be equivalently implemented in integrated circuits, as one or more computer programs running on one or more computers (e.g., as one or more programs running on one or more computer systems), as one or more programs running on one or more processors (e.g., as one or more programs running on one or more microprocessors), as firmware, or as virtually any combination thereof, and that designing the circuitry and/or writing the code for the software and or firmware would be well within the skill of one of skill in the art in light of this disclosure.

[0380] In addition, those skilled in the art will appreciate that the mechanisms of the subject matter described herein are capable of being distributed as a program product in a variety of forms, and that an illustrative aspect of the subject matter described herein applies regardless of the particular type of signal bearing medium used to actually carry out the distribution. Examples of a signal bearing medium include, but are not limited to, the following: a recordable type medium such as a floppy disk, a hard disk drive, a Compact Disc (CD), a Digital Video Disk (DVD), a digital tape, a computer memory, etc.; and a transmission type medium such as a digital and/or an analog communication medium (e.g., a fiber

optic cable, a waveguide, a wired communications link, a wireless communication link (e.g., transmitter, receiver, transmission logic, reception logic, etc.).

[0381] In summary, numerous benefits have been described which result from employing the concepts described herein. The foregoing description of the one or more aspects has been presented for purposes of illustration and description. It is not intended to be exhaustive or limiting to the precise form disclosed. Modifications or

¹⁰ variations are possible in light of the above teachings. The one or more aspects were chosen and described in order to illustrate principles and practical application to thereby enable one of ordinary skill in the art to utilize the various aspects and with various modifications as are ¹⁵ suited to the particular use contemplated. It is intended

that the claims submitted herewith define the overall scope.

Example 1. A surgical instrument, comprising: a handle assembly comprising: a drive system; an electric motor mechanically coupled to the drive system; a controller electrically coupled to the electric motor; and at least one user-actuated input device electrically coupled to the controller; and an interchangeable shaft comprising cutting member, wherein the interchangeable shaft is configured to engage the drive system of the handle assembly; wherein the at least one user-actuated input device is configured to control a firing speed of the cutting member when the interchangeable shaft is engaged with the handle assembly.

Example 2. The surgical instrument of example 1, wherein the at least one user-actuated input device comprises at least two switches.

Example 3. The surgical instrument of example 2, wherein one of the at least two switches is a speed control switch that controls a firing speed of the cutting member.

Example 4. The surgical instrument of example 3, wherein the speed control switch comprises a rotarystyle switch, a slider-style switch, or a rocker-style switch.

Example 5. The surgical instrument of example 2, wherein one of the at least two switches is an activation control switch.

Example 6. The surgical instrument of example 5, wherein the activation control switch comprises a button-style switch.

Example 7. The surgical instrument of example claim 3, wherein the speed control switch comprises a variable resistor.

Example 8. The surgical instrument of example claim 3, wherein the variable resistor is coupled to an Analog to Digital (A/D) Converter.

Example 9. A method for operating a surgical instrument comprising: implementing, via a controller of the surgical instrument, an operating mode of the surgical instrument, wherein the operating mode

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comprises operating a component of the surgical device according to a first speed; receiving, via the controller, an electrical signal associated with an activation of at least one user-actuated input device electrically coupled to the controller; and overriding, via the controller, the operating mode of the surgical instrument upon receiving the electrical signal; and operating, via the controller, the component of the surgical device according to a second speed that is greater than the first speed.

Example 10. The method of example 9, wherein the operating mode is a first operating mode and the component is a first component, the method further comprising implementing, via the controller of the surgical instrument, a second operating mode of the surgical instrument, wherein the second operating mode comprises operating a second component of the surgical device.

Example 11. The method example 10, wherein the first component is a cutting member and the second ²⁰ component is an articulation member.

Example 12. The method of example 9, further comprising sensing a parameter, via at least one sensor of a surgical instrument, associated with a drive system of the surgical instrument.

Example 13. The method of example 12, wherein the operating mode of the surgical instrument is based on the parameter sensed by the at least one sensor.

Example 14. The method of claim 9, wherein operating the component of the surgical device according to the second speed comprises operating the component to return the component of the surgical device to a base position.

Example 15. The method of claim 9, wherein overriding the operating mode comprises completing an operation associated the component of the surgical device prior to operating the component of the surgical device according to the second speed.

Example 16. A surgical instrument, comprising: a drive system; an electric motor mechanically coupled to the drive system; a controller electrically coupled to the electric motor, wherein the controller is configured to implement an operating mode of the surgical instrument, wherein the controller implementing the operating mode comprises the controller operating a component of the surgical device according to a first speed; at least one user-actuated input device electrically coupled to the controller; wherein the controller is configured to receive an electrical signal associated with an activation of the at least one user-actuated input device, to override the operating mode of the electric motor upon receiving the electrical signal, and to operate the component of the surgical device according to a second speed that is greater than the first speed upon overriding the operating mode.

Example 17. The surgical instrument of example 16,

wherein the operating mode is a first operating mode and the component is a first component, and wherein the controller is configured to implement a second operating mode of the surgical instrument upon completing an operation of the first component, and wherein the second operating mode comprises operating a second component of the surgical device. Example 18. The surgical instrument example 17, wherein the first component is a cutting member and the second component is an articulation member.

Example 19. The surgical instrument of example 16, further comprising at least one sensor configured to sense a parameter associated with the surgical instrument.

Example 20. The surgical instrument of example 19, wherein the parameter associated with the surgical instrument is associated with the drive system of the surgical instrument, and wherein the operating mode of the surgical instrument is based on the parameter sensed by the at least one sensor.

Example 21. The surgical instrument of example 16, wherein the parameter associated with the surgical instrument is a tissue parameter associated with tissue engaged by the surgical instrument.

Example 22. The surgical instrument of example 16, wherein the first component is a cutting member and wherein the controller is configured to override the first operating mode of the electric motor upon receiving the electrical signal and to operate the cutting member to return the cutting member to a base position.

Example 23. A surgical instrument, comprising: a drive system; an end effector comprising an anvil, an elongated channel mechanically coupled to the anvil, a staple cartridge supported by the elongated channel and a cutting member mechanically coupled to the drive system; an electric motor mechanically coupled to the drive system; at least one sensor electrically coupled to the controller and configured to sense a tissue parameter associated with tissue engaged with the end effector; a controller electrically coupled to the electric motor and electrically coupled to the sensor, wherein the controller is configured to implement an operating mode of the surgical instrument based on the tissue parameter, and wherein the controller implementing the operating mode comprises the controller operating a first component of the surgical device; and at least one user-actuated input device electrically coupled to the controller; and wherein the controller is configured to receive an electrical signal associated with an activation of the at least one user-actuated input device; and wherein, upon receiving the electrical signal, the controller is configured to complete an operation associated with the operating mode of the surgical instrument; and wherein, upon completion of the operation associated with the operating mode, the controller is configured to return the first component of the surgical de-

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vice to a base position; and wherein, upon returning the first component to the base position, the controller is configured to operate a second component of the surgical device.

Example 24. The surgical instrument of example 23, wherein the operating mode is a first operating mode, and wherein the controller is configured to implement a second operating mode of the surgical instrument upon returning the first component to the base position.

Example 25. The surgical instrument example 24, wherein the first component is a cutting member and the second component is an articulation member.

Example 26. The surgical instrument of example 24, wherein the controller is configured to operate the first component of the surgical device at a first speed and the controller is configured to return the first component of the surgical device to the base position at a second speed, wherein the second speed is greater than the first speed.

Example 27. The surgical instrument of example 24, wherein the at least sensor comprises a first sensor and a second sensor, wherein the first sensor is configured to sense the tissue parameter and wherein the second sensor is configured to drive system parameter associated with the drive system of the surgical instrument, and wherein the controller is configured to implement the first operating mode of the surgical instrument based on the tissue parameter and the drive system parameter.

Example 28. The surgical instrument of example 24, further comprising a non-transitory computer readable medium, and wherein the non-transitory computer readable medium stores instructions associated with the operating mode.

[0382] The following is a non-exhaustive list of numbered embodiments which may be claimed:

- 1. A surgical instrument, comprising:
 - a drive system;

an electric motor mechanically coupled to the drive system;

a controller electrically coupled to the electric ⁴⁵ motor, and

at least one user-actuated input device electrically coupled to the controller; and

wherein the controller is configured to operate the electric motor according to an operating ⁵⁰ mode for the electric motor;

wherein the controller is configured to detect activation of the at least one user-actuated input device and to override the operating mode for the electric motor upon detecting the activation of the at least one user-actuated input device.

2. The surgical instrument of embodiment 1, further

comprising a sensor electrically coupled to the controller, wherein the sensor is configured to sense a parameter associated with the drive system, and wherein the controller is configured to implement the operating mode for the electric motor based on the parameter sensed by the sensor.

3. The surgical instrument of embodiment 2, wherein the controller is configured to adjust power to the electric motor if the parameter exceeds a predefined threshold value according to operating mode for the electric motor.

4. The surgical instrument of embodiment 1, further comprising an end effector comprising an anvil and a cutting member mechanically coupled to the drive system.

5. The surgical instrument of embodiment 4, further comprising a second sensor configured to sense a second parameter associated with the cutting member.

6. The surgical instrument of embodiment 1, further comprising a battery electrically coupled to the electric motor.

7. The surgical instrument of embodiment 1, wherein the controller is configured to operate the electric motor according to an input received via the at least one user-actuated input device upon the controller overriding the operating system of electric motor.

8. The surgical instrument of embodiment 1, wherein the controller is configured to operate the electric motor according to an input associated with the at least one user-actuated input device upon the controller overriding the operating system of electric motor and when the parameter sensed by the sensor meets or exceeds a predefined threshold value.

9. The surgical instrument of embodiment 1, further comprising a user selection display electrically coupled to the controller.

10. The surgical instrument of embodiment 9, wherein the controller is configured to receive, via an input from the user selection display electrically, a command that configures the controller to override the operating mode for the electric motor upon receiving the activation of the at least one user-actuated input device.

11. The surgical instrument of embodiment 2, wherein the controller is configured to implement a predetermined operation of the electric motor when the parameter sensed by the sensor meets or exceeds a predefined threshold value according to the oper-

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ating mode for the electric motor.

12. A surgical instrument, comprising:

a drive system;

an end effector comprising an anvil, an elongated channel mechanically coupled to the anvil, a staple cartridge supported by the elongated channel and a cutting member mechanically coupled to the drive system;

an electric motor mechanically coupled to the drive system;

a controller electrically connected to the electric motor;

a first sensor electrically coupled to the controller and configured to sense a first parameter associated with the anvil;

a second sensor electrically coupled to the controller and configured to sense a second parameter associated with the cutting member; and at least one user-actuated input device electri-

cally coupled to the controller; and

wherein the controller is configured to implement an operating mode for the electric motor based on at least one of the first parameter and the second parameter; and

wherein the controller is configured such that activation of the at least one user-actuated input device causes the controller to override the operating mode for the electric motor.

13. The surgical instrument of embodiment 12, wherein the controller is configured to adjust an input to the electric motor if at least one of the first parameter or the second parameter exceeds a predefined ³⁵ threshold value according to the operating mode for the electric motor.

14. The surgical instrument of embodiment 12, further comprising a battery electrically coupled to the electric motor.

15. The surgical instrument of embodiment 12, wherein the controller is configured to operate the electric motor according to an input received via the at least one user-actuated input device upon the controller overriding the operating system of electric motor.

16. The surgical instrument of embodiment 12, ⁵⁰ wherein the controller is configured to operate the electric motor according to an input associated with the at least one user-actuated input device upon the controller overriding the operating system of electric motor. ⁵⁵

17. A method for operating a surgical instrument comprising:

sensing a parameter, via a sensor of a surgical instrument, associated with a drive system of the surgical instrument;

implementing, via a controller of the surgical instrument, an operating mode of an electric motor of the surgical instrument;

receiving, via the controller, an electrical signal; and

overriding, via the controller, the operating mode of the electric motor upon receiving the electrical signal; and

wherein the electrical signal comprises an activation of at least one user-actuated input device electrically coupled to the controller.

18. The method of embodiment 17, wherein the operating mode of the surgical instrument is based on the parameter sensed by the sensor.

19. The method of embodiment 17, wherein implementing the operating mode of the electric motor comprises the controller adjusting power to the electric motor if the parameter exceeds a predefined threshold value.

20. The method of embodiment 17, further comprising receiving, via an input from a user selection display of the surgical instrument, a command that configures the controller to override the operating mode for the electric motor upon receiving the activation of the at least one user-actuated input device.

21. The method of embodiment 17, further comprising operating the electric motor according to an input received via the at least one user-actuated input device upon the controller overriding the operating system of electric motor.

Claims

1. A surgical instrument, comprising:

a drive system;

an electric motor mechanically coupled to the drive system;

a controller electrically coupled to the electric motor, and

at least one user-actuated input device electrically coupled to the controller; and

wherein the controller is configured to operate the electric motor according to an operating mode for the electric motor;

wherein the controller is configured to detect activation of the at least one user-actuated input device and to override the operating mode for the electric motor upon detecting the activation of the at least one user-actuated input device.

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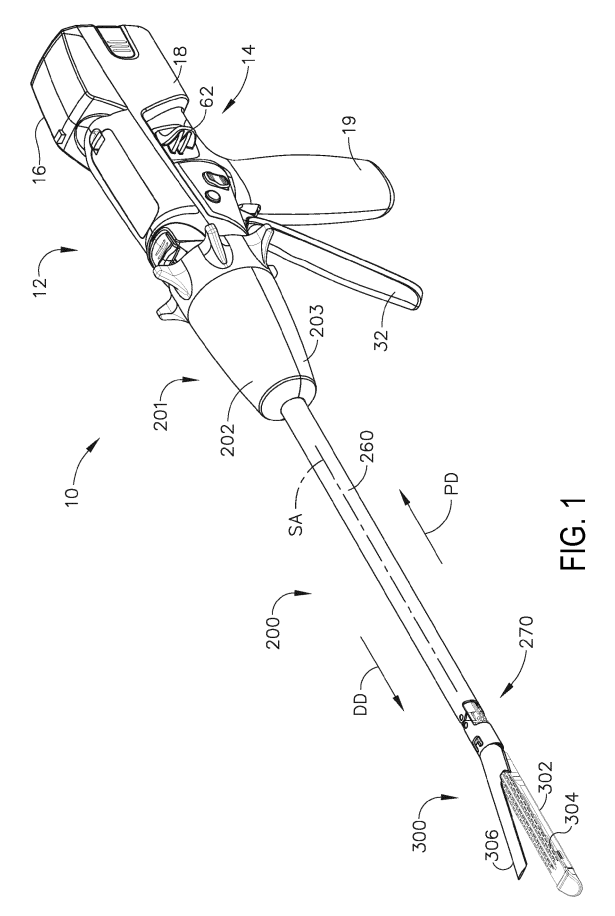
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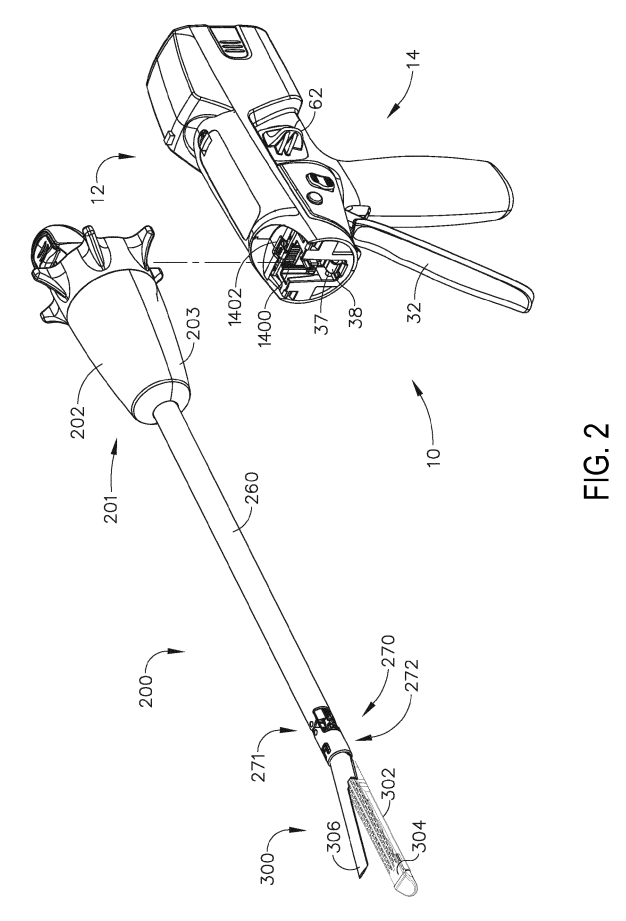
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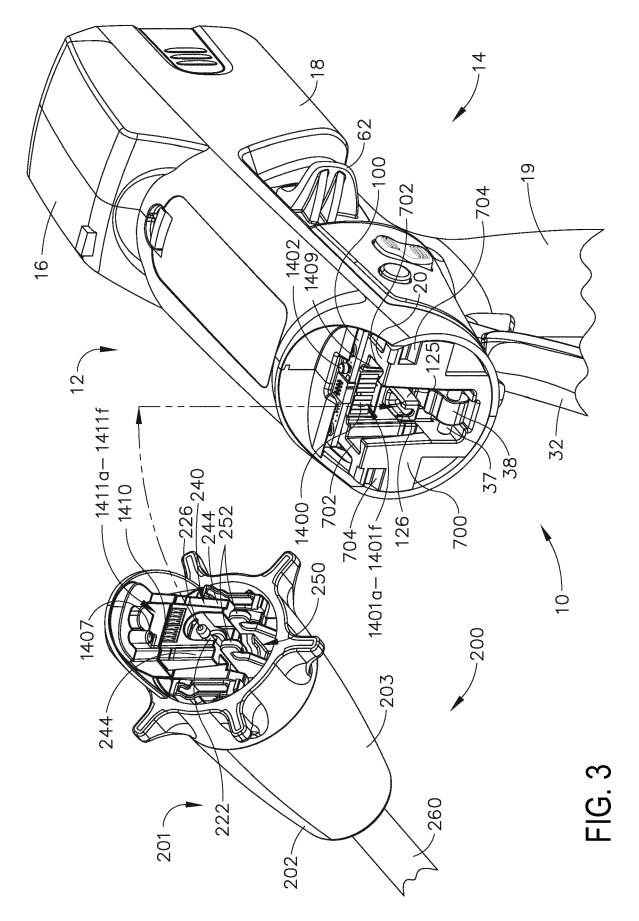
- 2. The surgical instrument of claim 1, further comprising a sensor electrically coupled to the controller, wherein the sensor is configured to sense a parameter associated with the drive system, and wherein the controller is configured to implement the operating mode for the electric motor based on the parameter sensed by the sensor.
- **3.** The surgical instrument of claim 2, wherein the controller is configured to adjust power to the electric ¹⁰ motor if the parameter exceeds a predefined threshold value according to operating mode for the electric motor.
- The surgical instrument of any one of the preceding claims, further comprising an end effector comprising an anvil and a cutting member mechanically coupled to the drive system.
- **5.** The surgical instrument of claim 4, further comprising ²⁰ a second sensor configured to sense a second parameter associated with the cutting member.
- The surgical instrument of any one of the preceding claims, further comprising a battery electrically coupled to the electric motor.
- 7. The surgical instrument of any one of the preceding claims, wherein the controller is configured to operate the electric motor according to an input received via the at least one user-actuated input device upon the controller overriding the operating system of electric motor.
- 8. The surgical instrument of any one of claims 1 to 6, ³⁵ wherein the controller is configured to operate the electric motor according to an input associated with the at least one user-actuated input device upon the controller overriding the operating system of electric motor and when the parameter sensed by the sensor ⁴⁰ meets or exceeds a predefined threshold value.
- **9.** The surgical instrument of any one of the preceding claims, further comprising a user selection display electrically coupled to the controller.
- 10. The surgical instrument of claim 9, wherein the controller is configured to receive, via an input from the user selection display electrically, a command that configures the controller to override the operating mode for the electric motor upon receiving the activation of the at least one user-actuated input device.
- The surgical instrument of any one of claims 2 to 10, wherein the controller is configured to implement a predetermined operation of the electric motor when the parameter sensed by the sensor meets or exceeds a predefined threshold value according to the

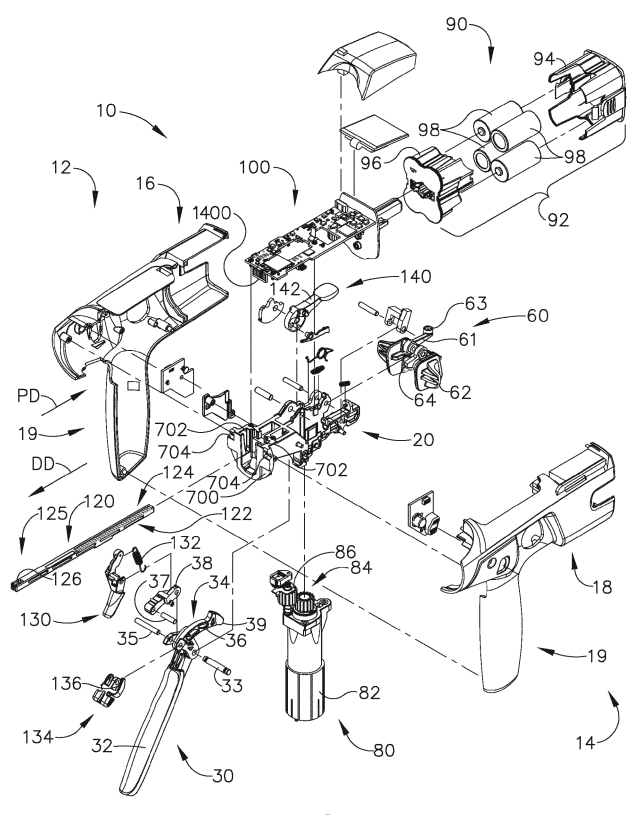
operating mode for the electric motor.

- **12.** A surgical instrument, comprising:
 - a drive system; an end effector comprising an anvil, an elongated channel mechanically coupled to the anvil, a staple cartridge supported by the elongated channel and a cutting member mechanically coupled to the drive system;
 - an electric motor mechanically coupled to the drive system;
 - a controller electrically connected to the electric motor;
 - a first sensor electrically coupled to the controller and configured to sense a first parameter associated with the anvil;
 - a second sensor electrically coupled to the controller and configured to sense a second parameter associated with the cutting member; and
 - at least one user-actuated input device electrically coupled to the controller; and
 - wherein the controller is configured to implement an operating mode for the electric motor based on at least one of the first parameter and the second parameter; and
 - wherein the controller is configured such that activation of the at least one user-actuated input device causes the controller to override the operating mode for the electric motor.
- **13.** The surgical instrument of claim 12, wherein the controller is configured to adjust an input to the electric motor if at least one of the first parameter or the second parameter exceeds a predefined threshold value according to the operating mode for the electric motor.
- **14.** The surgical instrument of claim 12 or claim 13, further comprising a battery electrically coupled to the electric motor.
- **15.** The surgical instrument of any one of claims 12 to 14, wherein the controller is configured to operate the electric motor according to one of: an input received via the at least one user-actuated input device upon the controller overriding the operating system of electric motor; and an input associated with the at least one user-actuated input device upon the controller overriding system of electric motor.

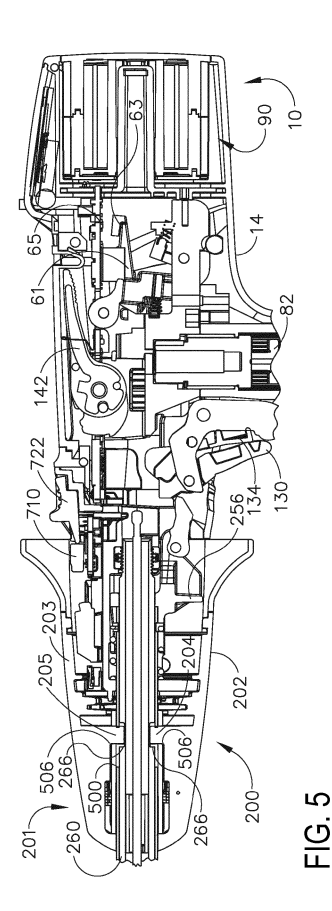


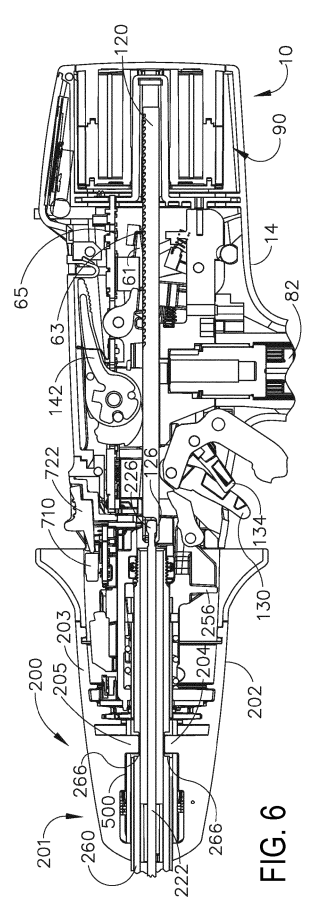


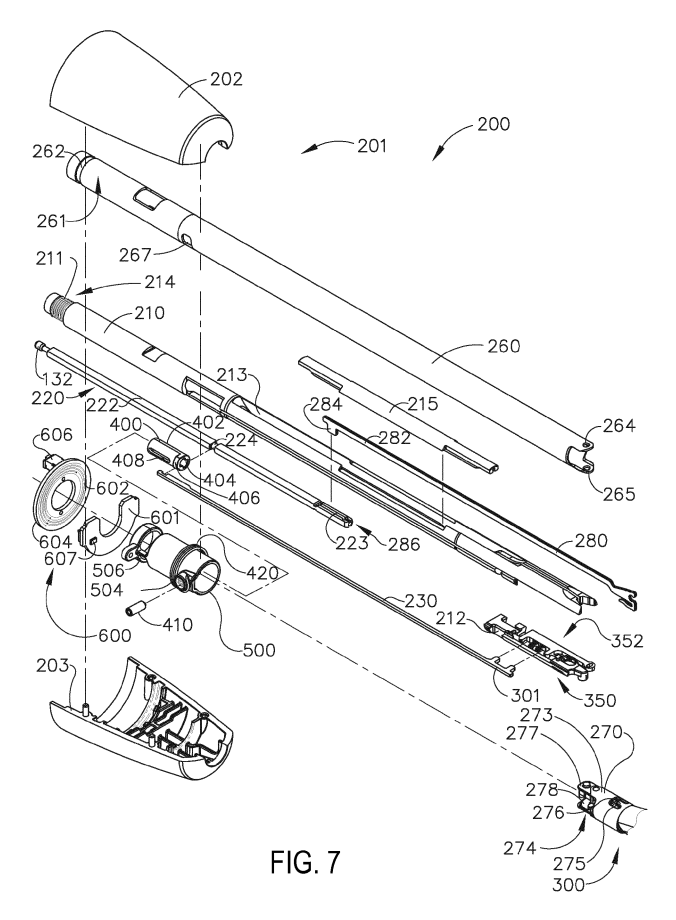


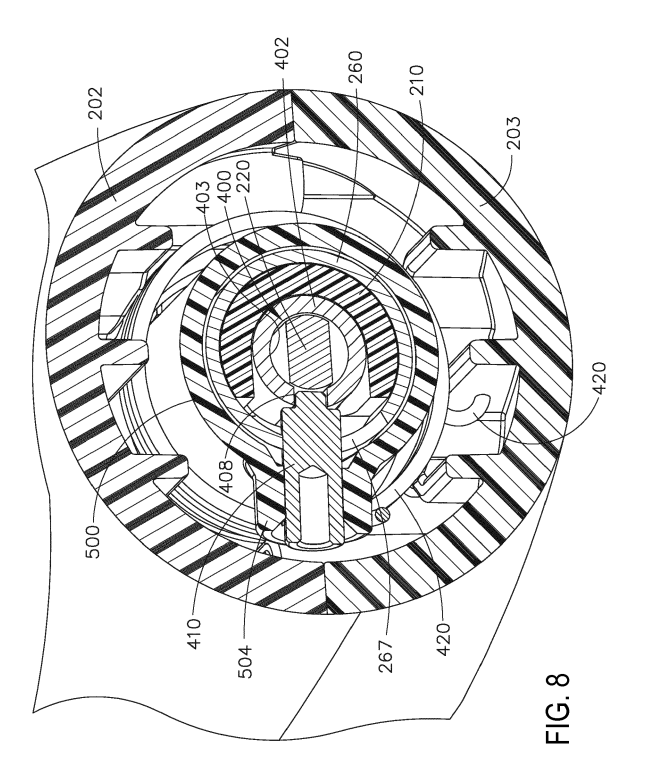


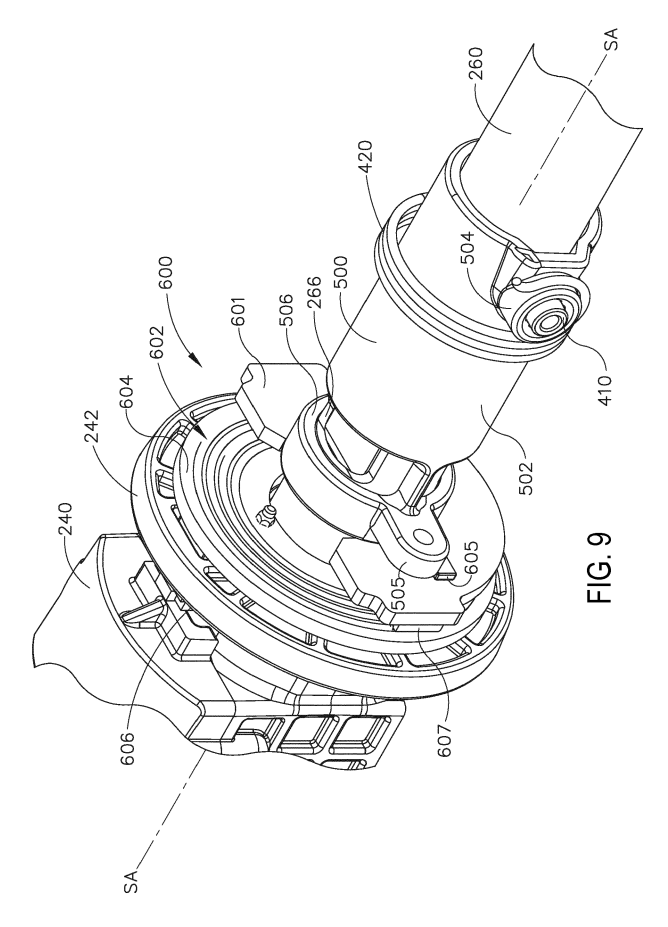


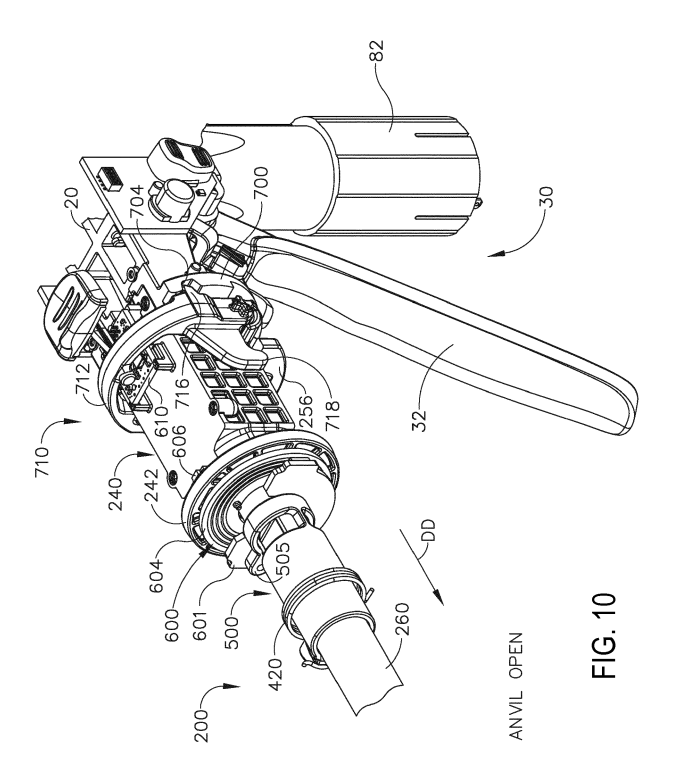


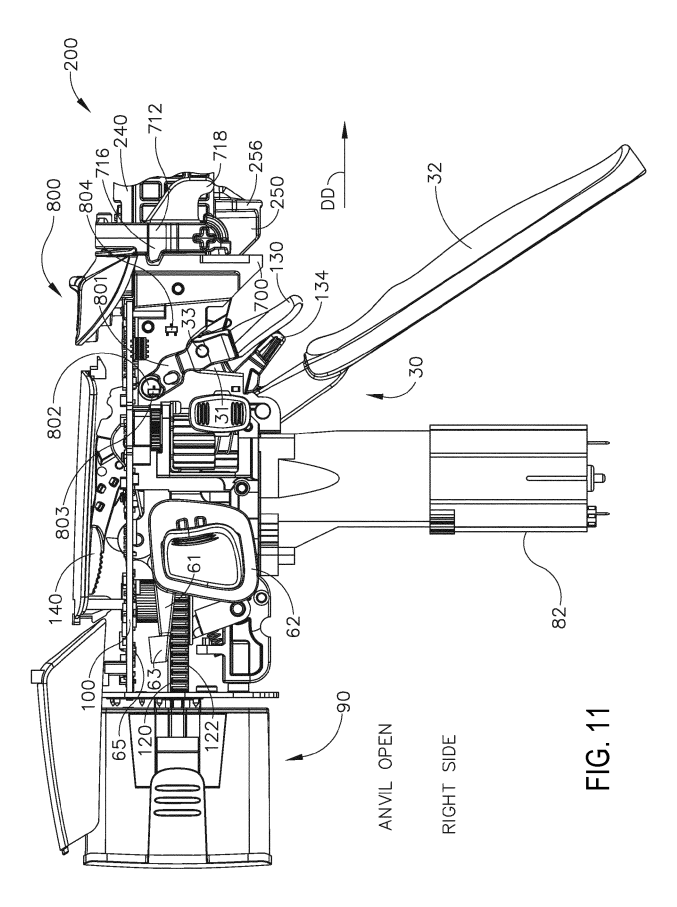


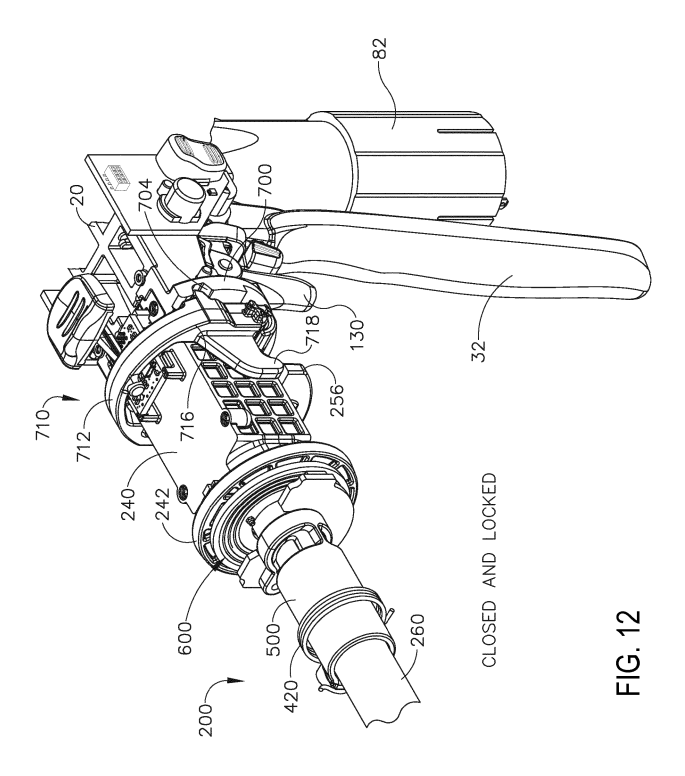


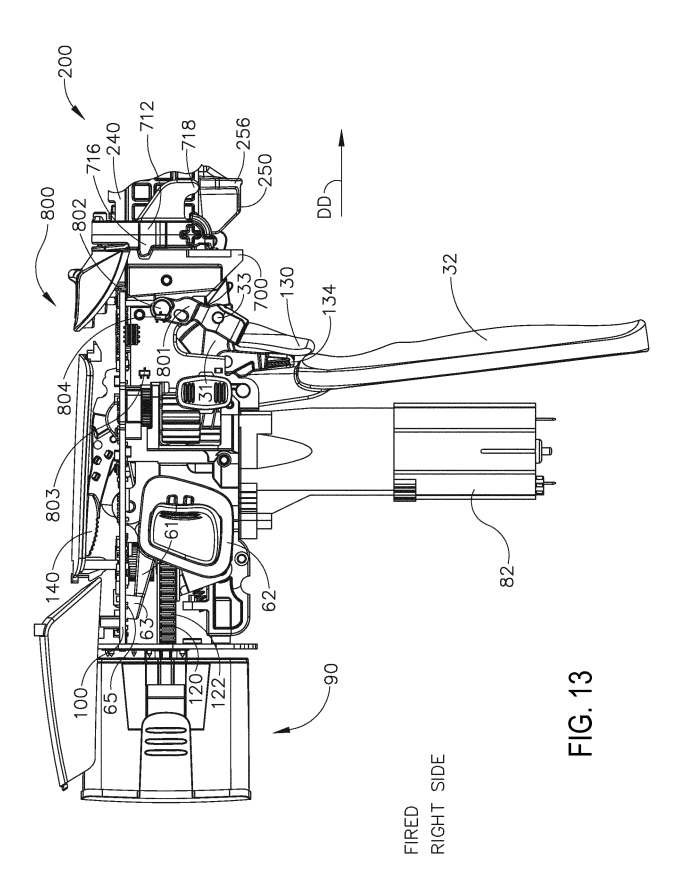












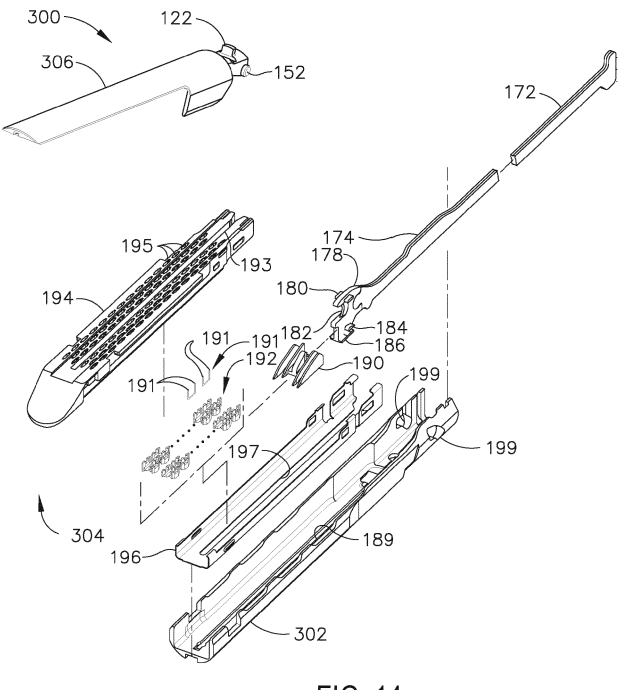
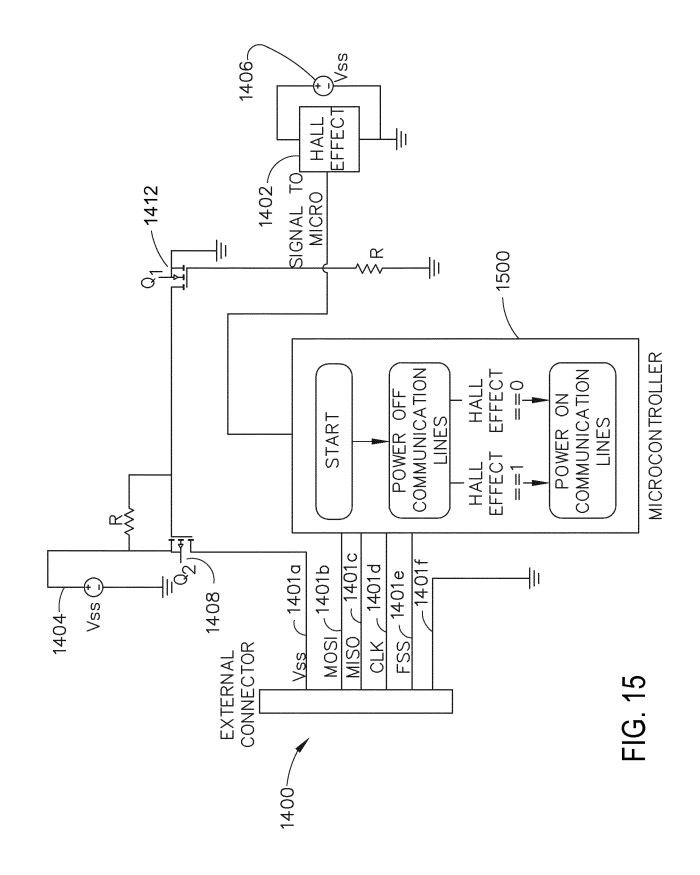
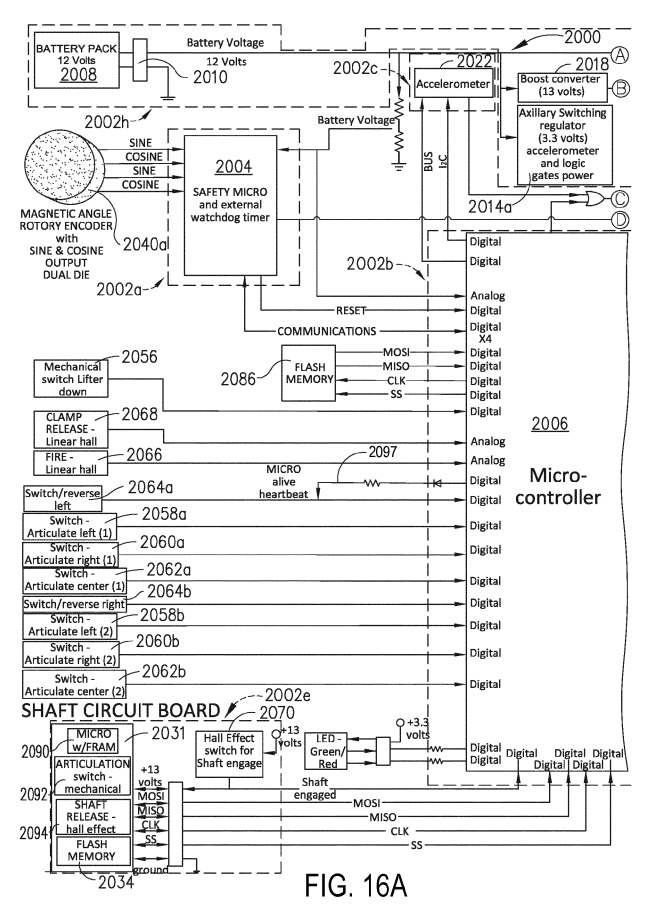
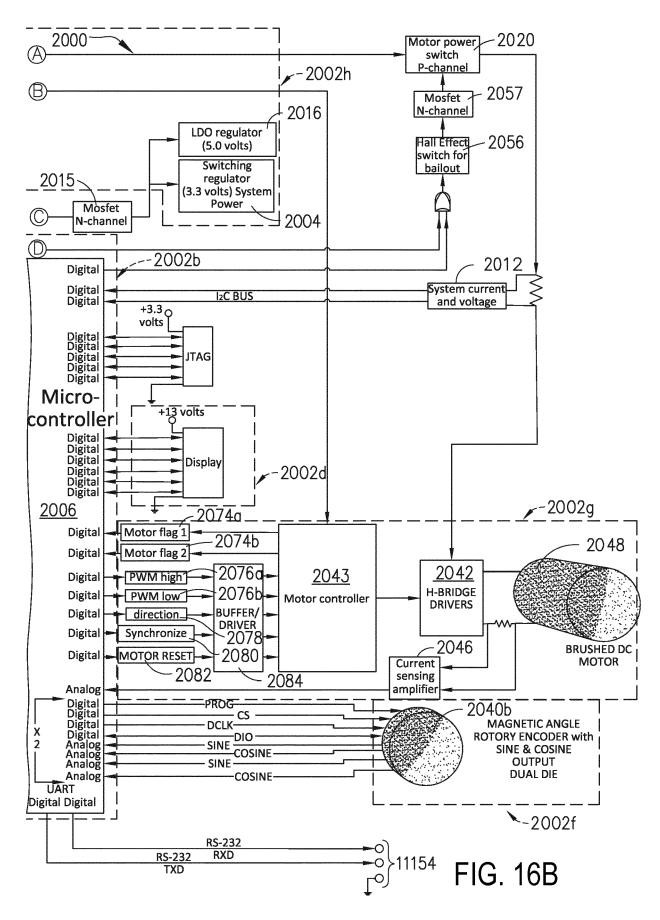


FIG. 14







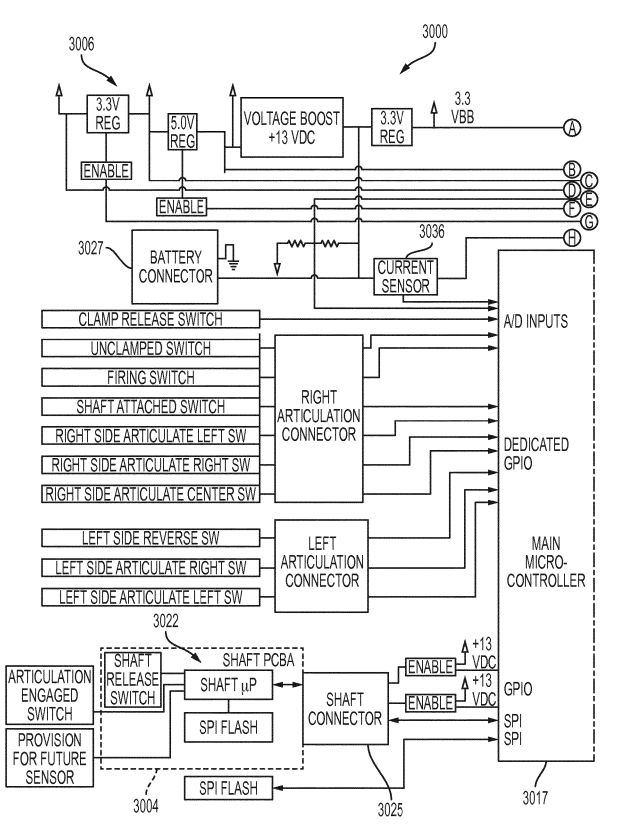


FIG. 17A

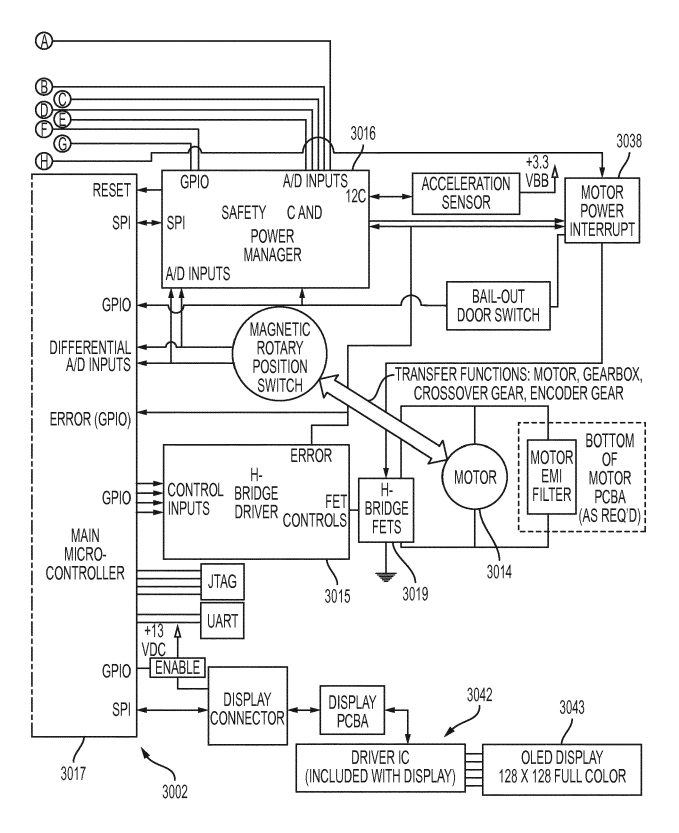
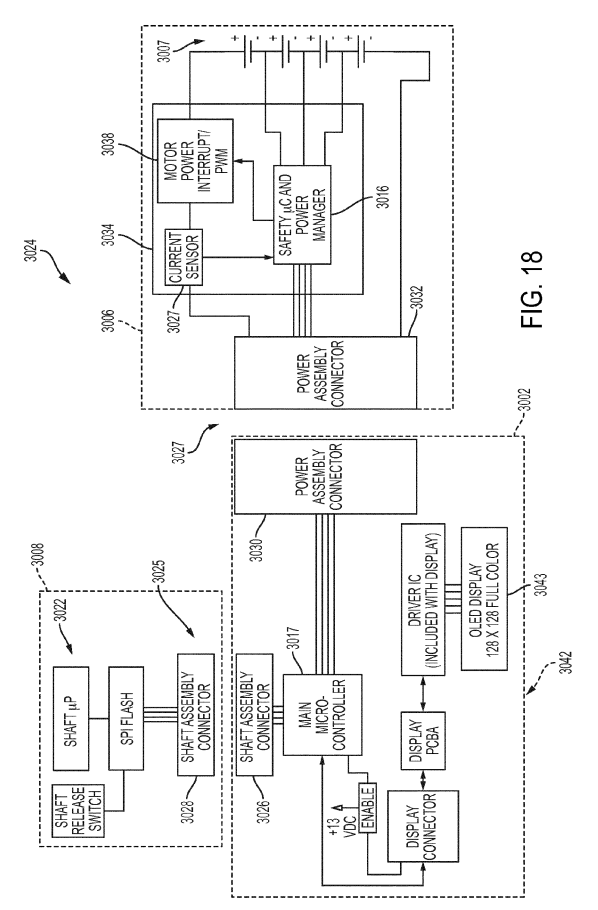
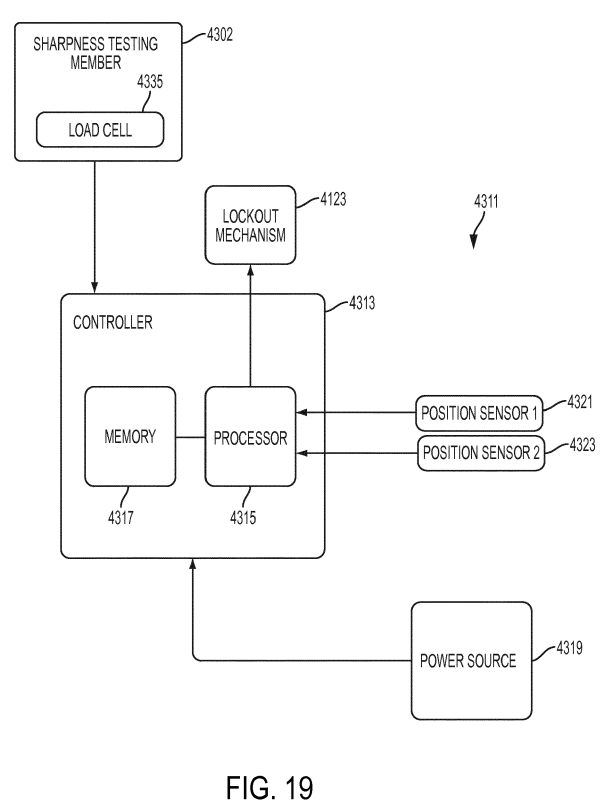


FIG. 17B

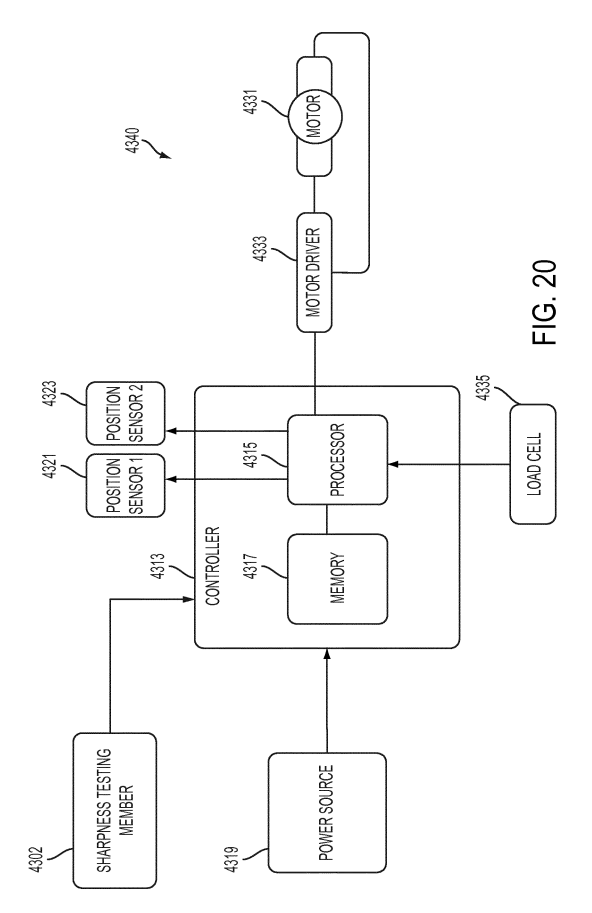
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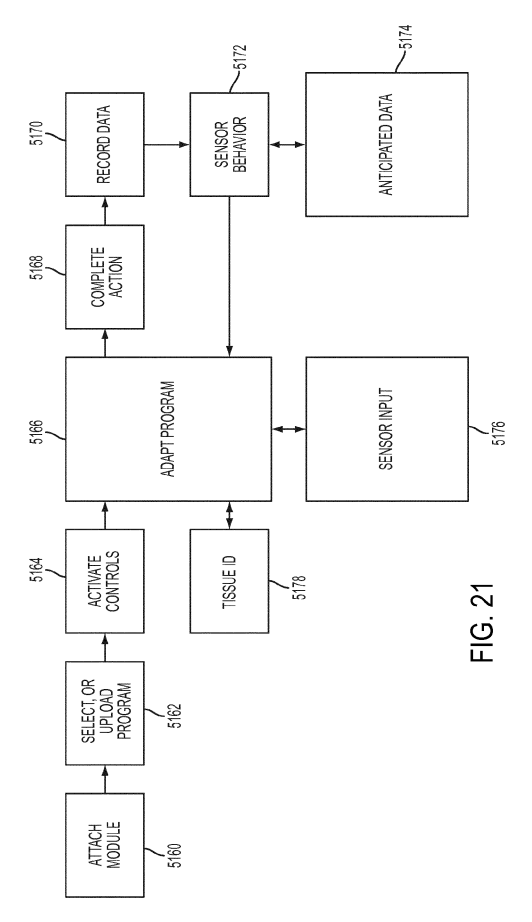


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10.13





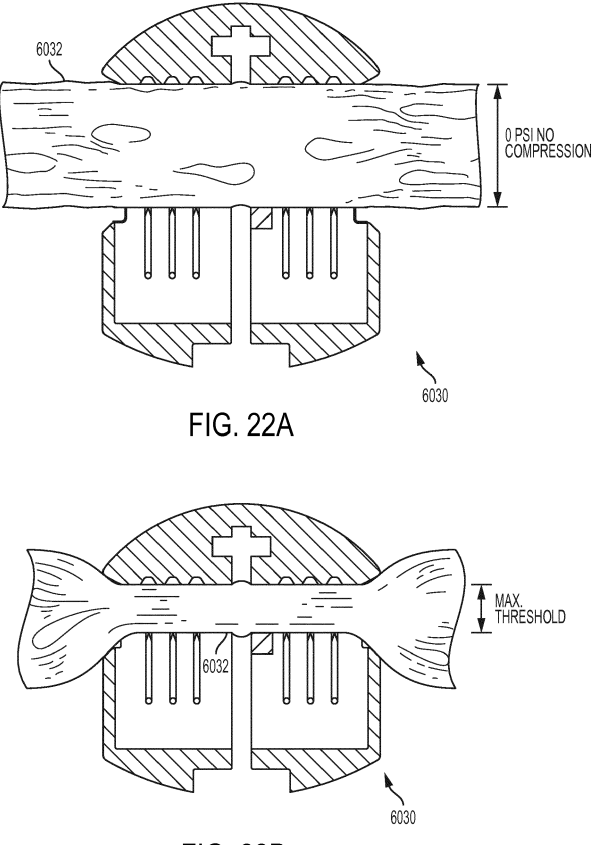


FIG. 22B

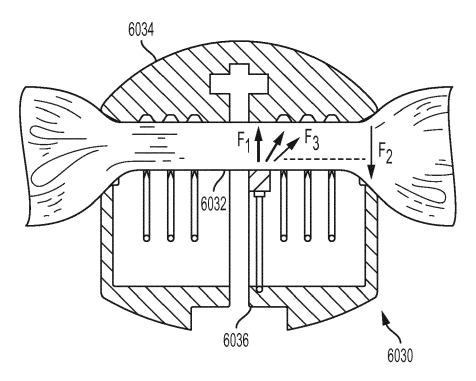


FIG. 23A

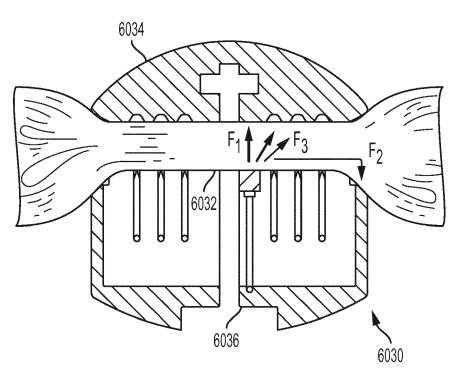
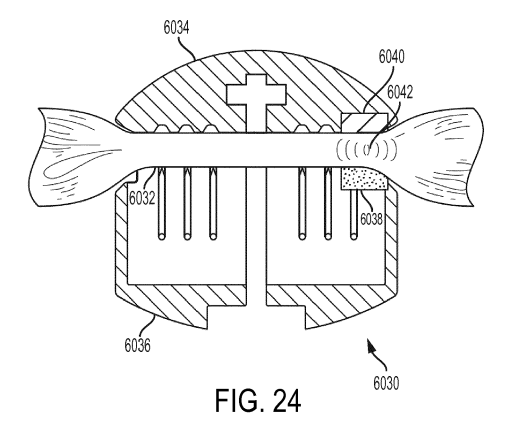
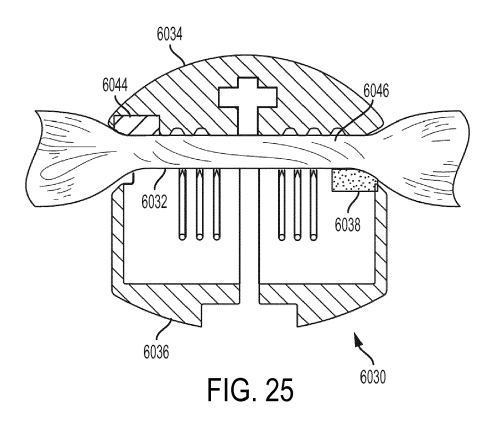
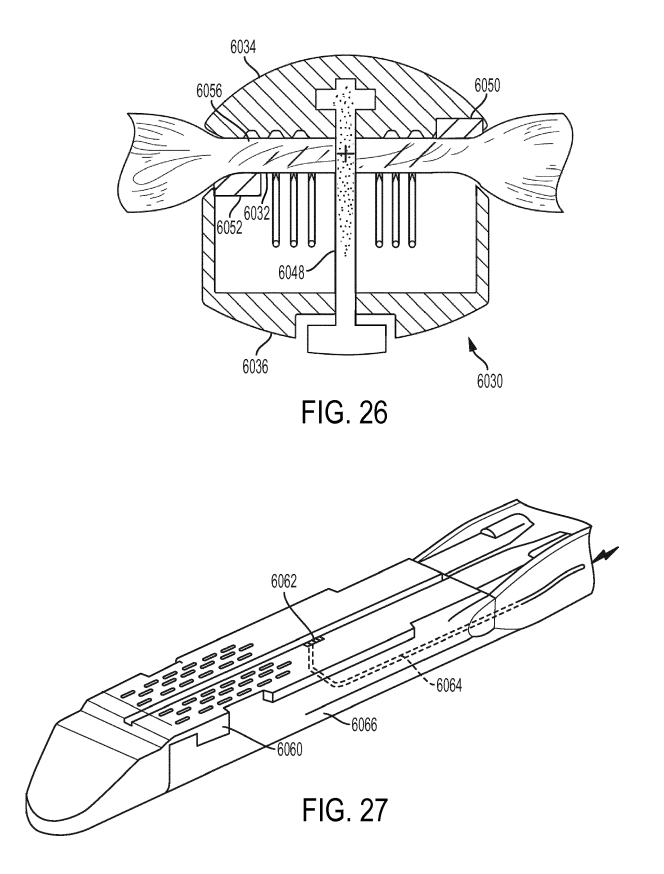


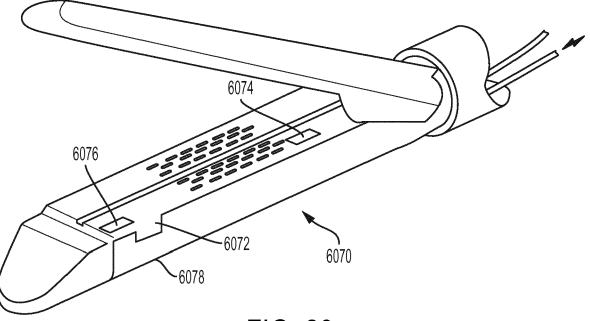
FIG. 23B



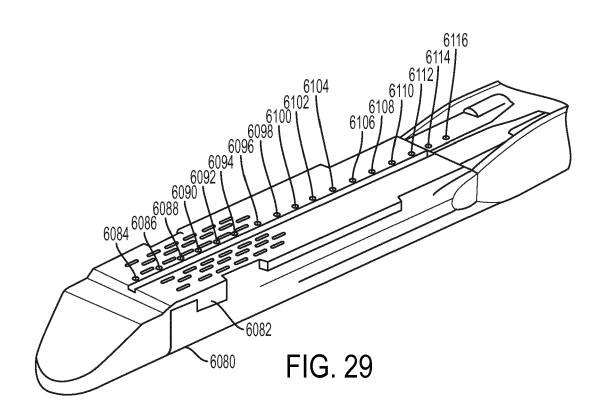


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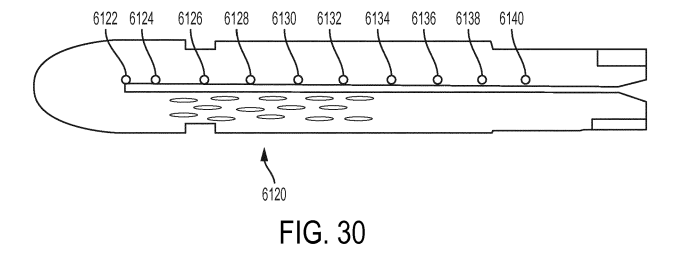


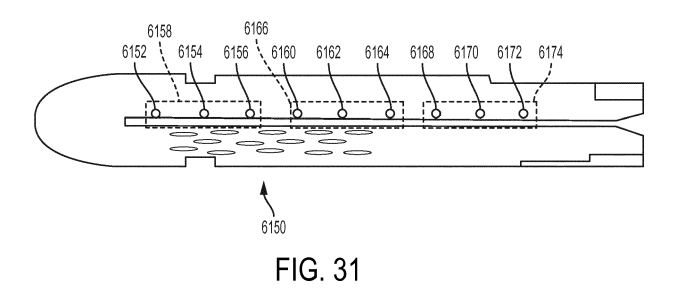


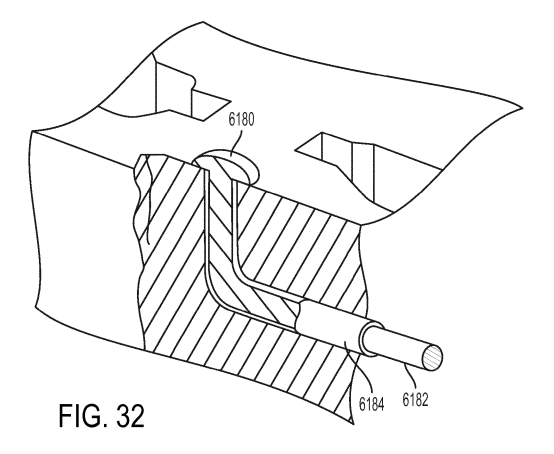












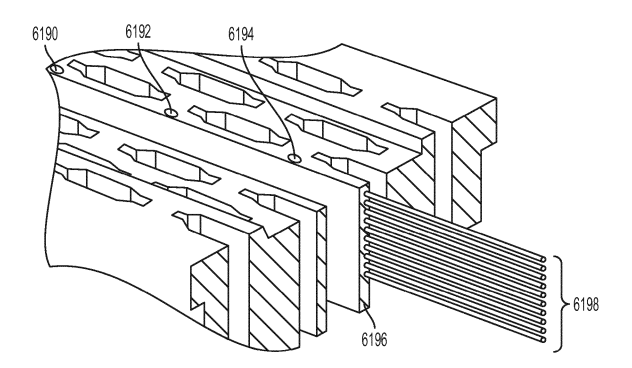
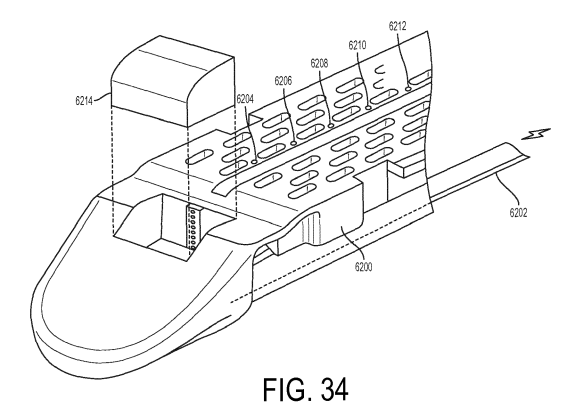
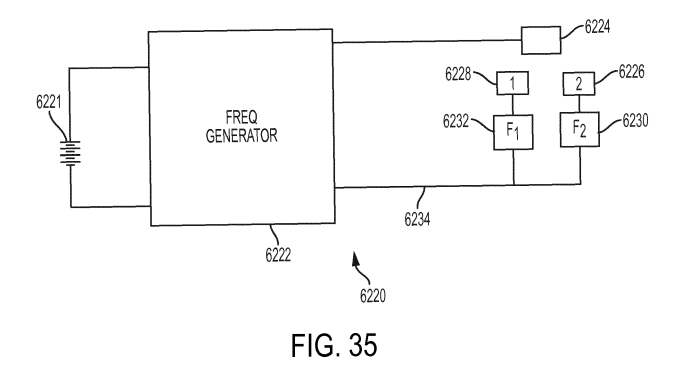


FIG. 33





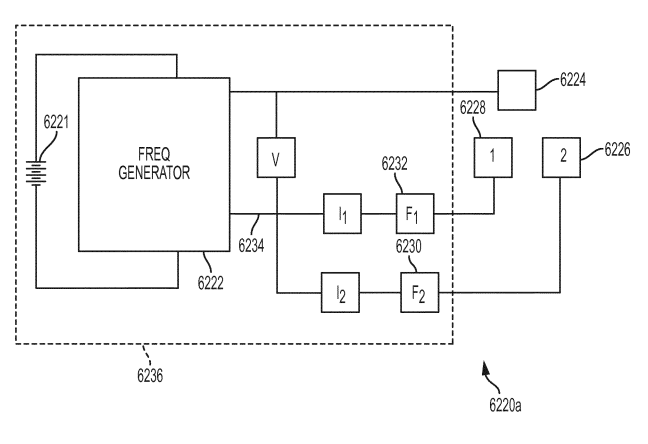
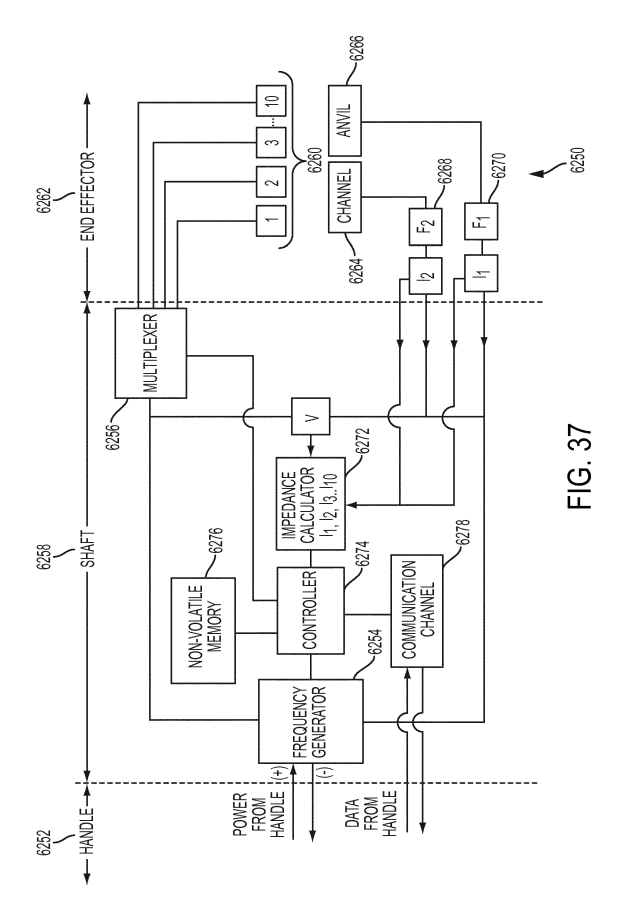
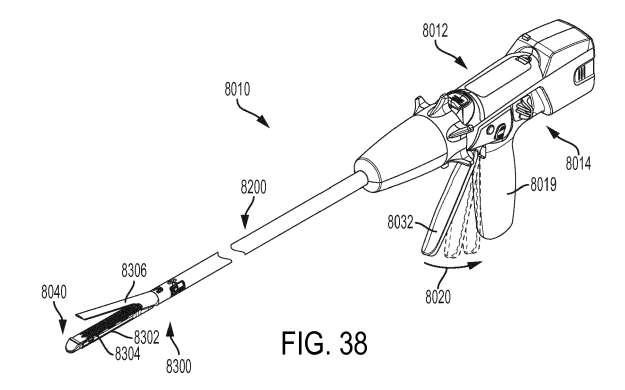
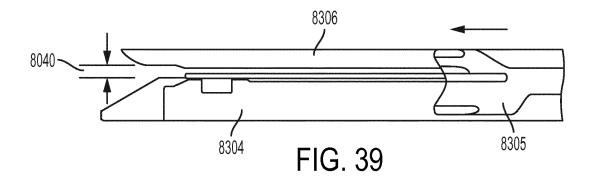
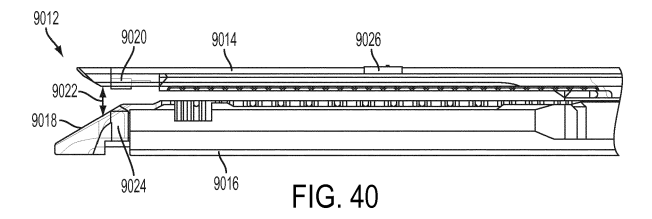


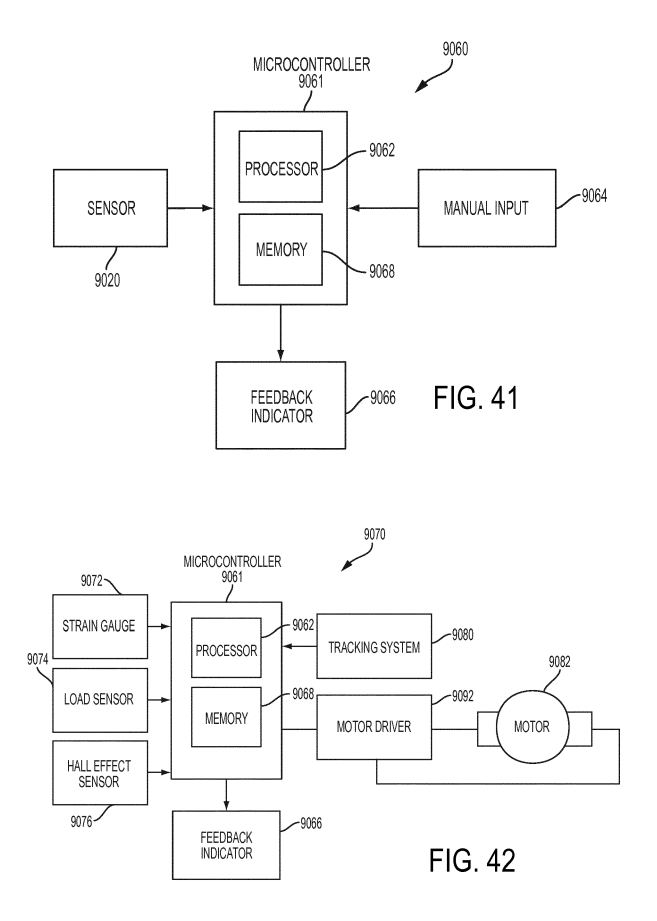
FIG. 36

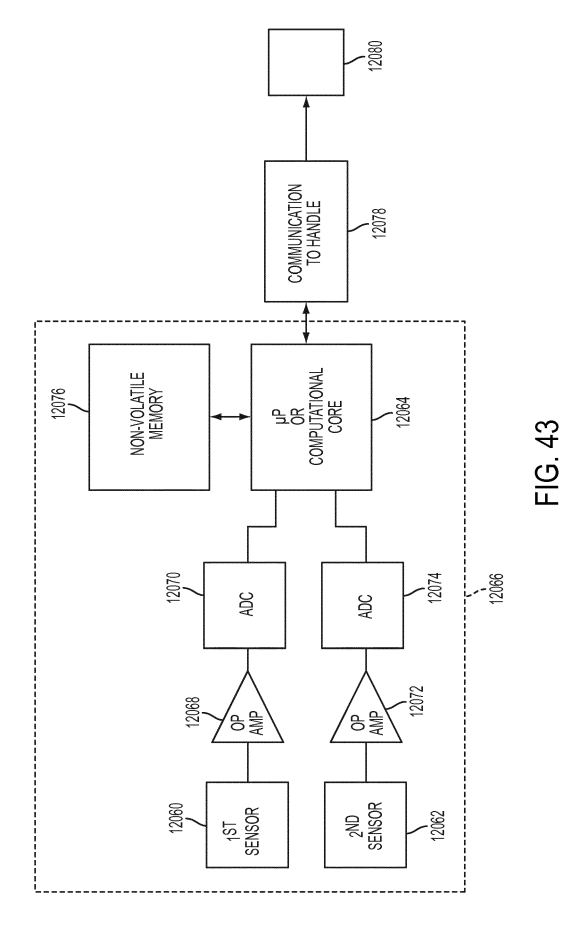


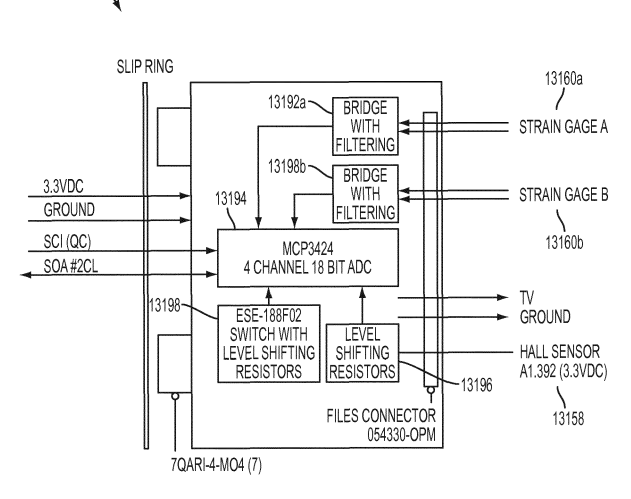








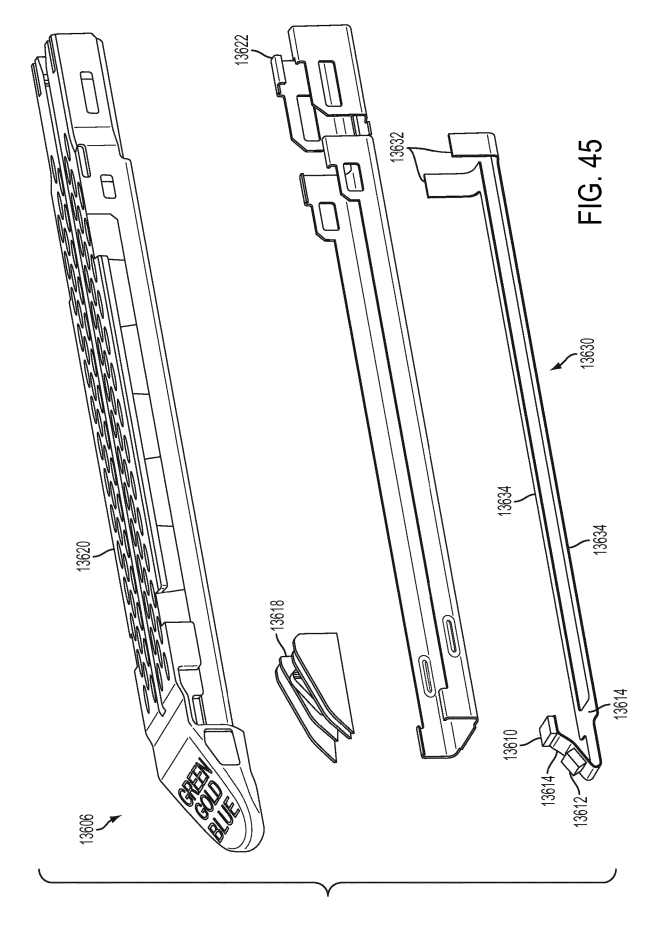


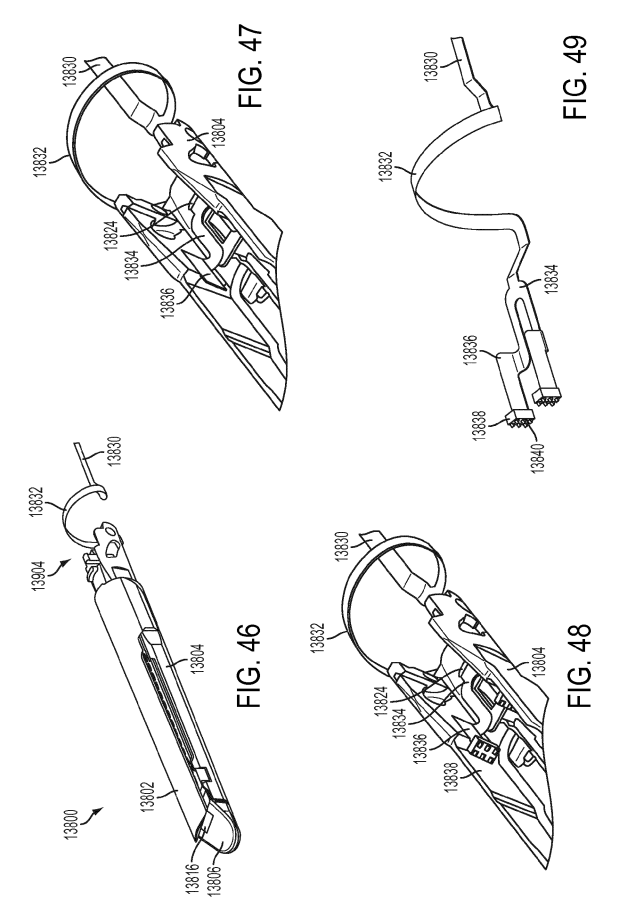


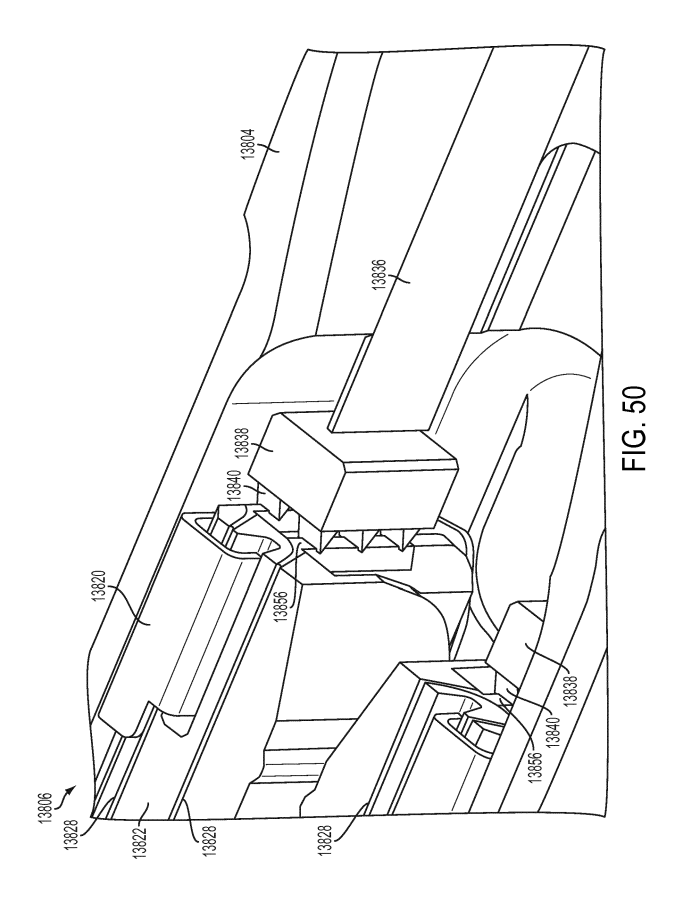
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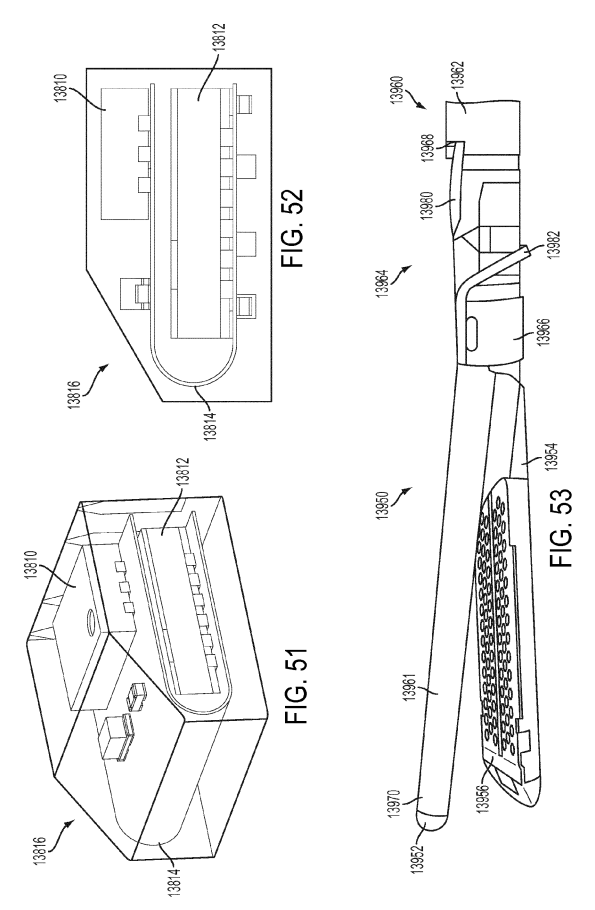
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FIG. 44









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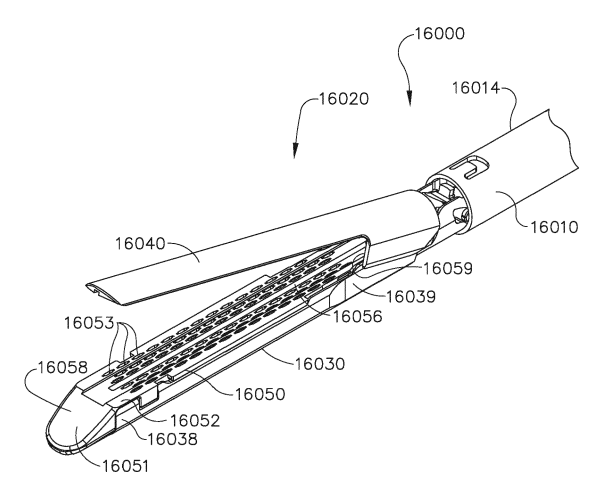
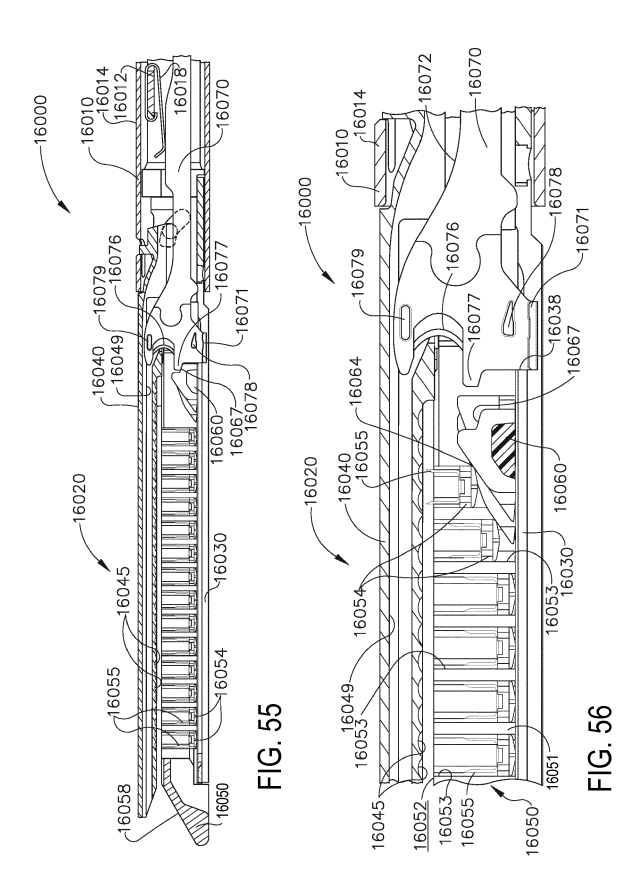
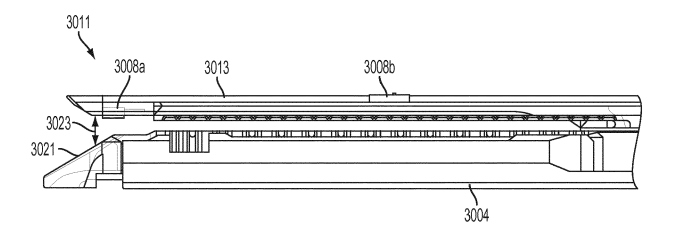
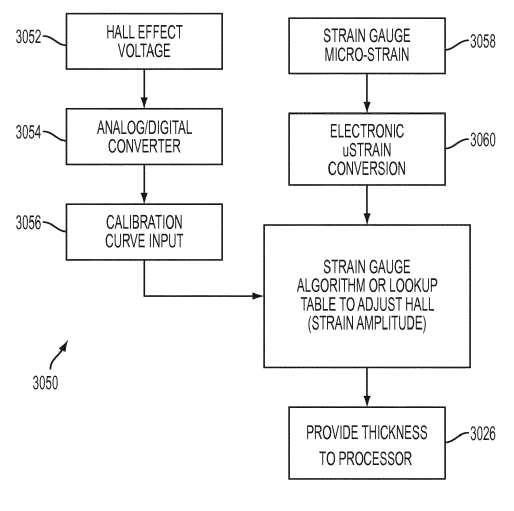


FIG. 54



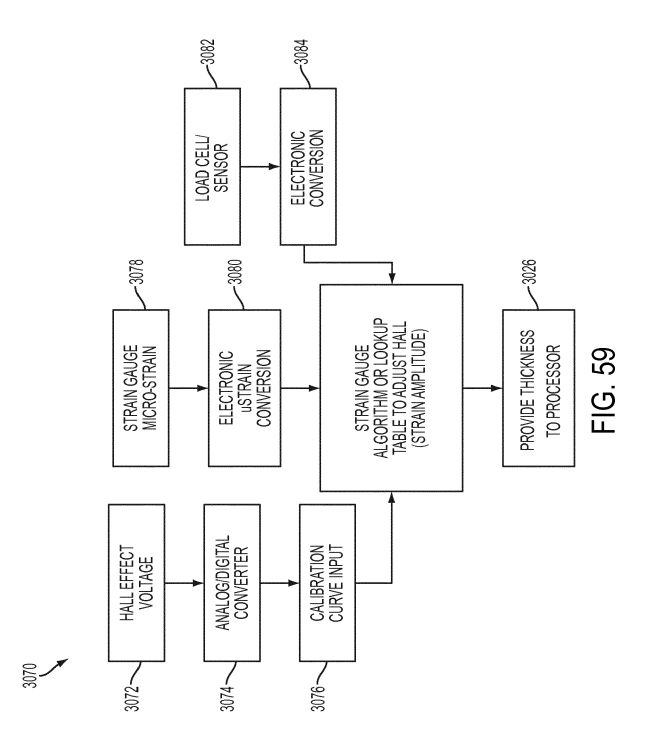


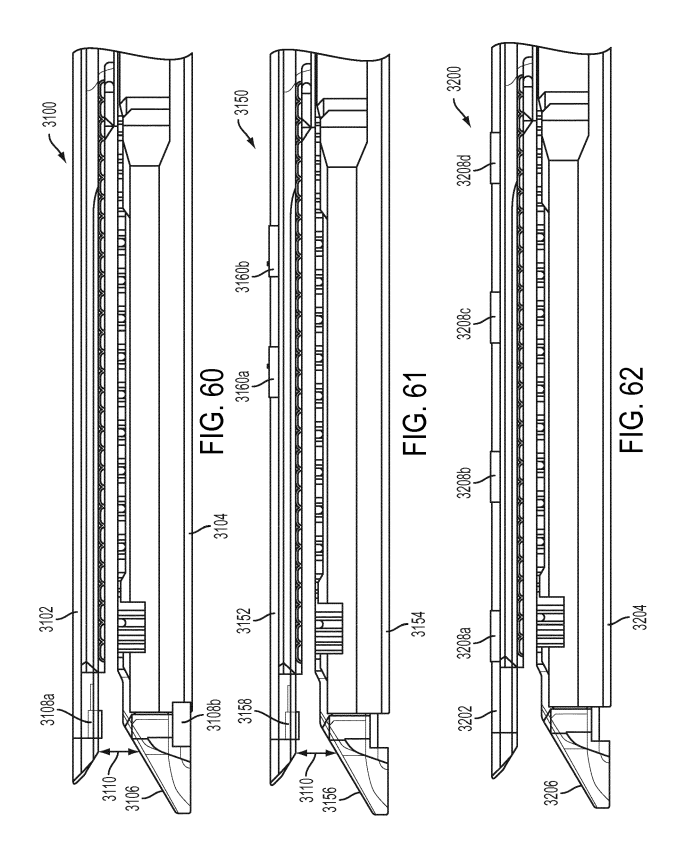


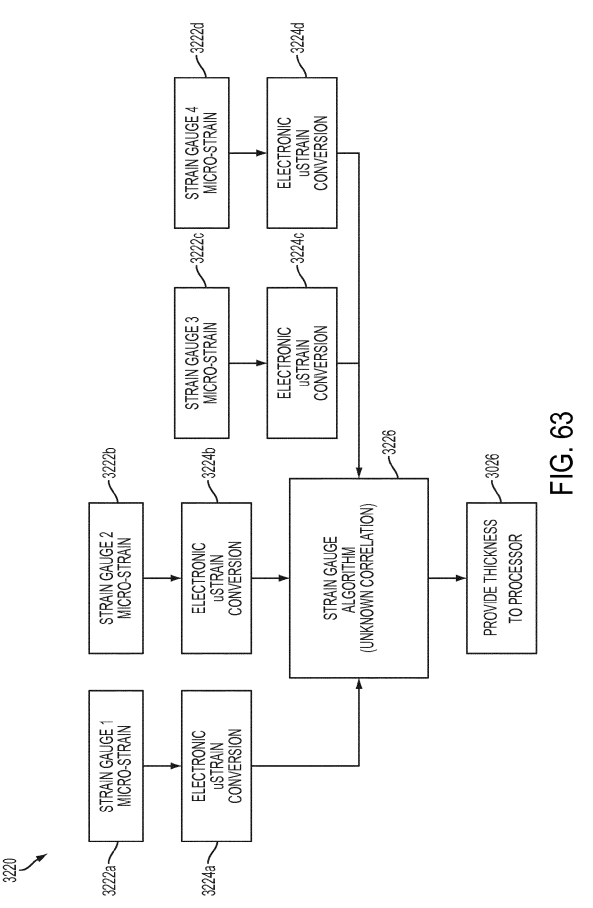




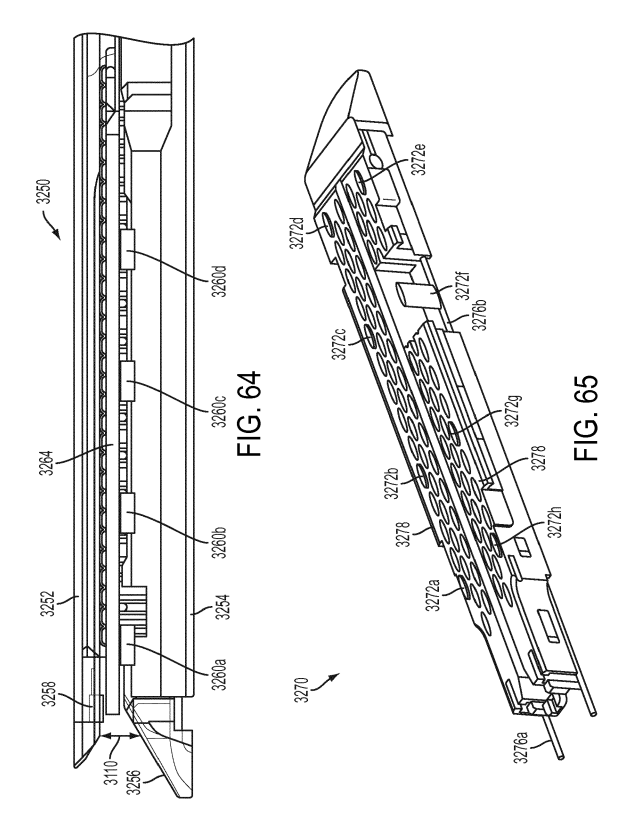


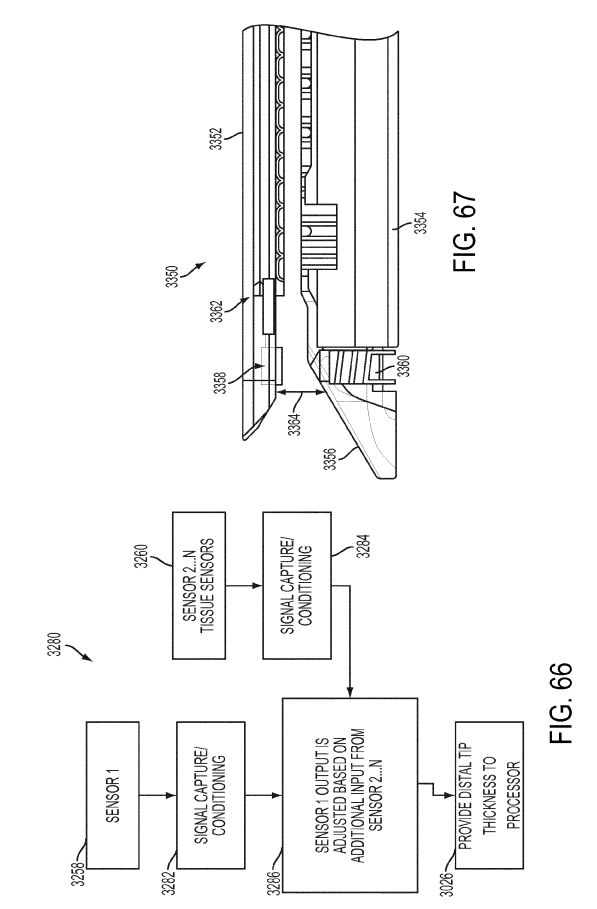


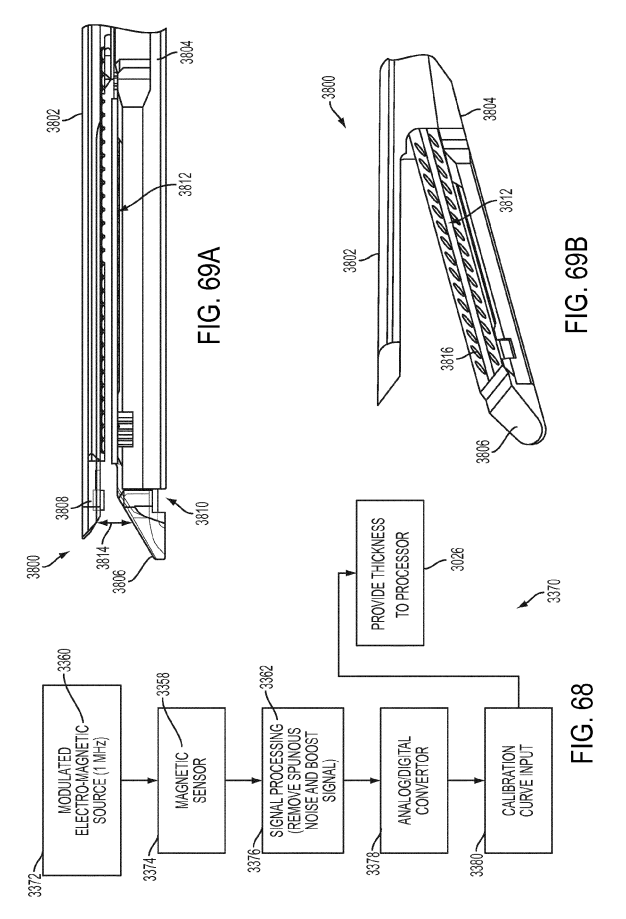


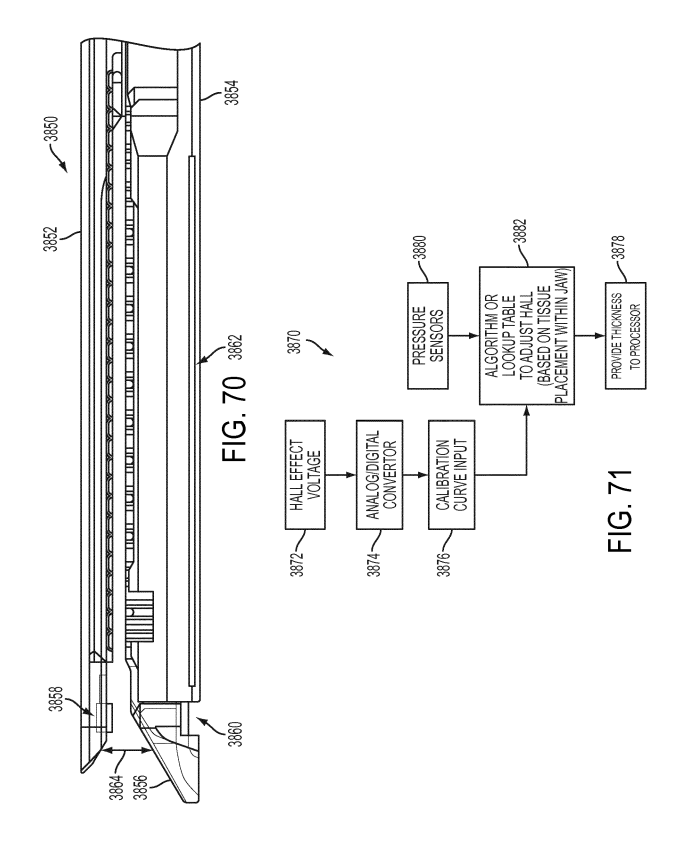


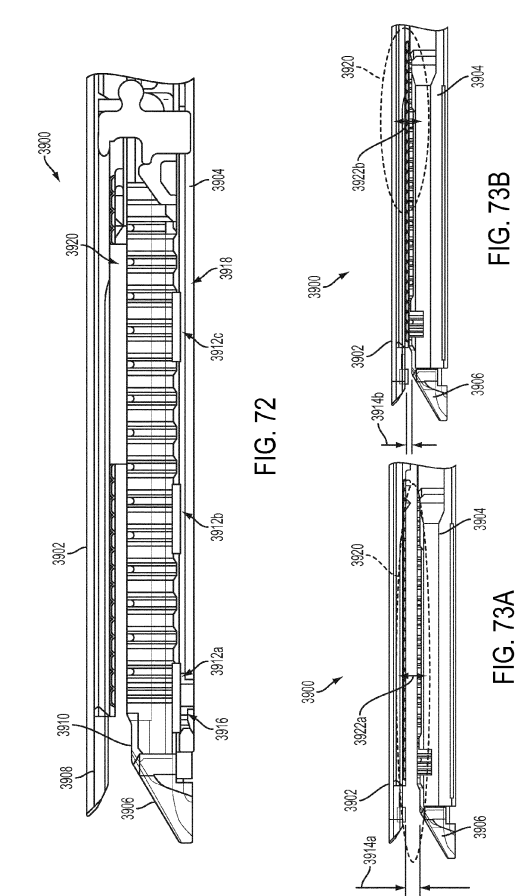
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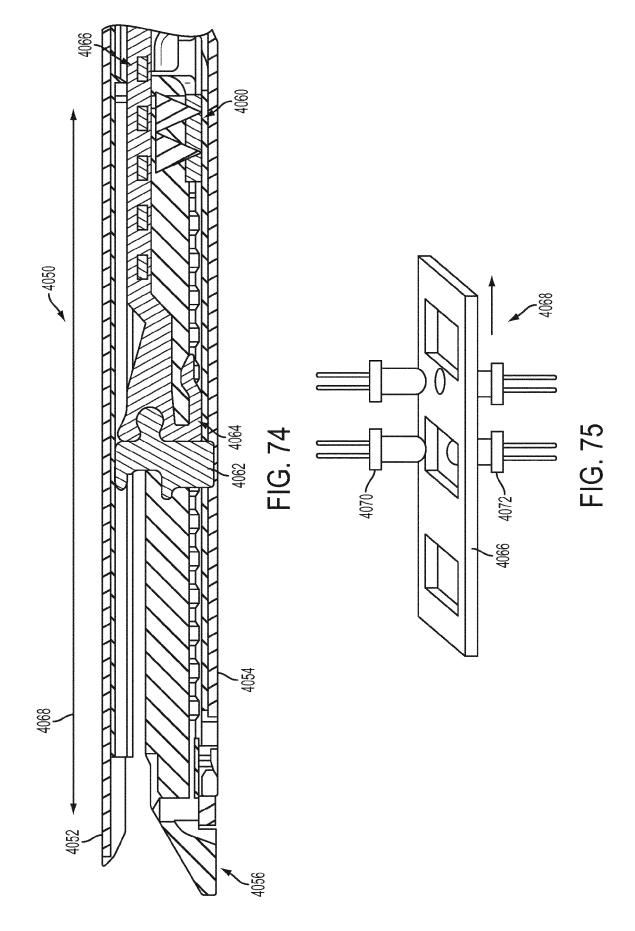


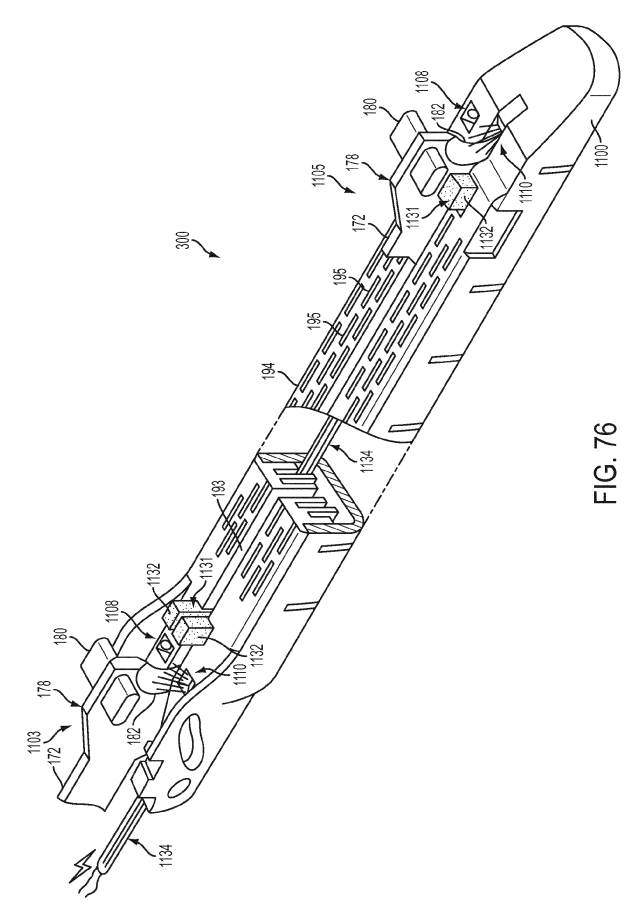


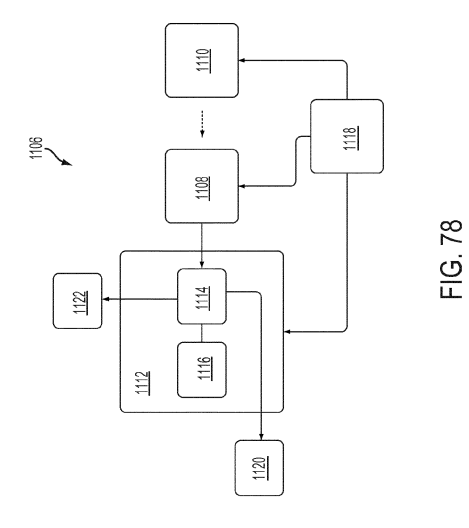


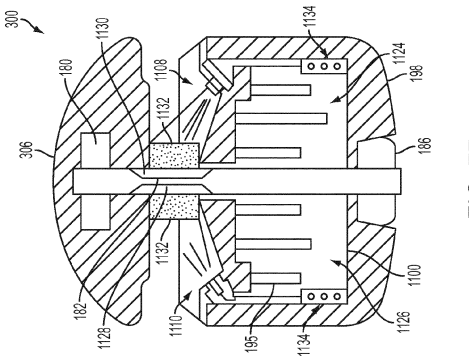


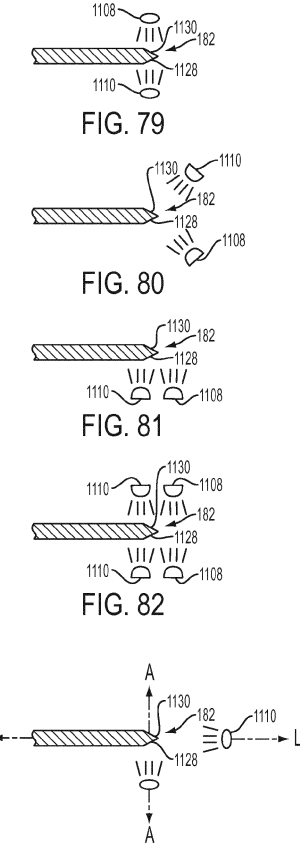




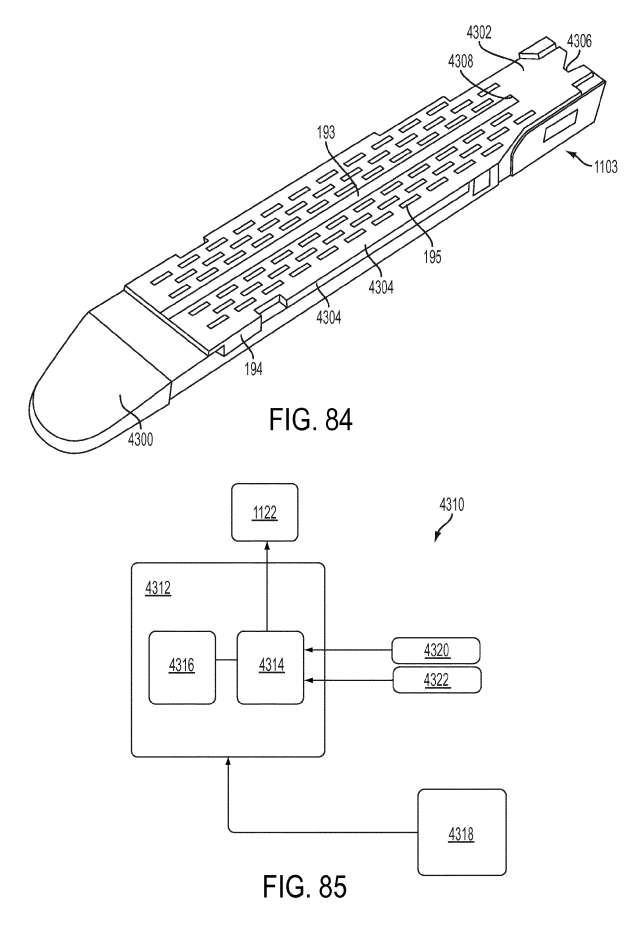


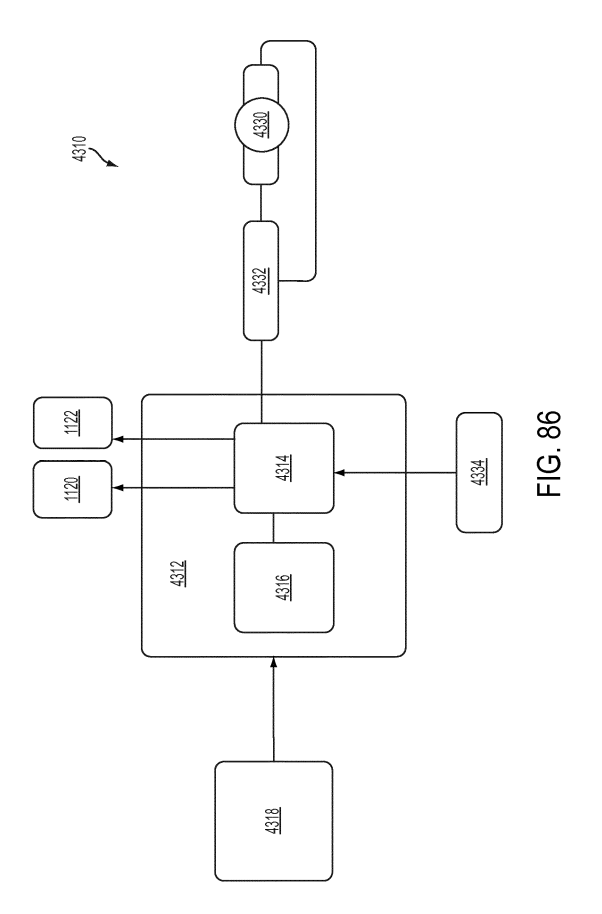


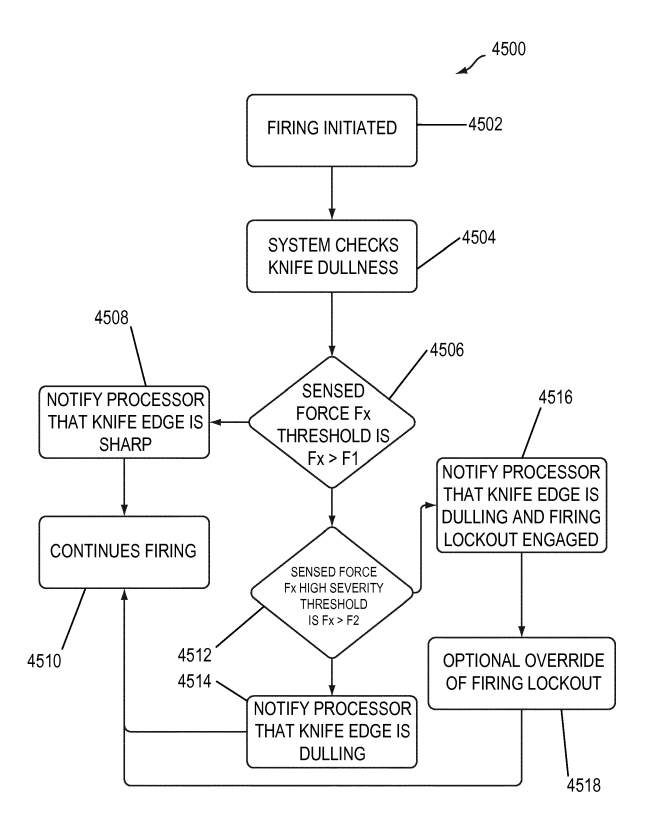


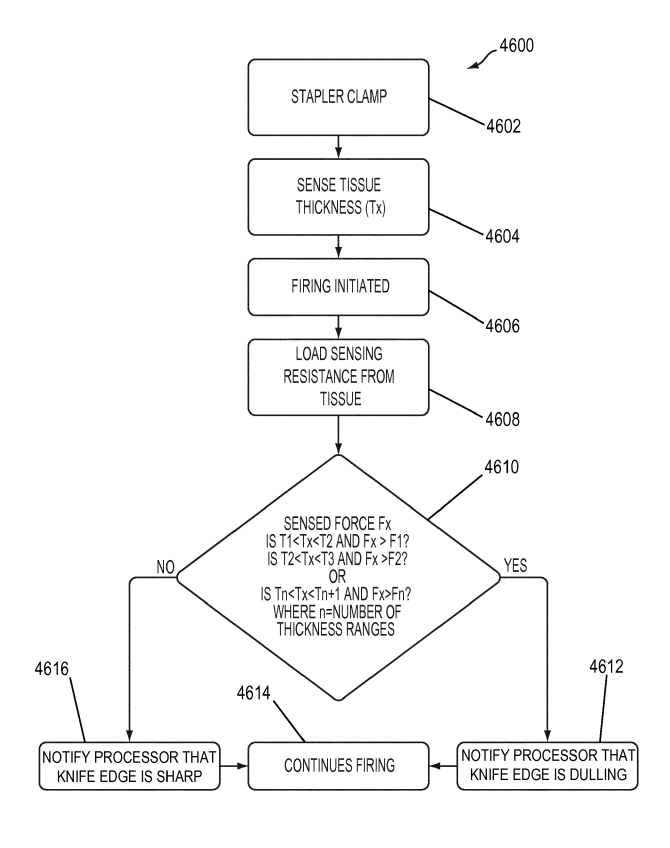






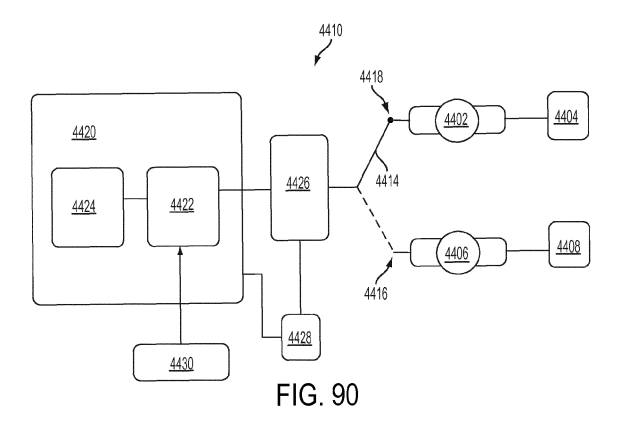








TISSUE THICKNESS RANGES (T)	THRESHOLD FORCE (F)
T1-T2	F1
т2-т3	F2
Т3-Т4	F3
-	-
-	-
-	-
Tn-1 - Tn	Fn



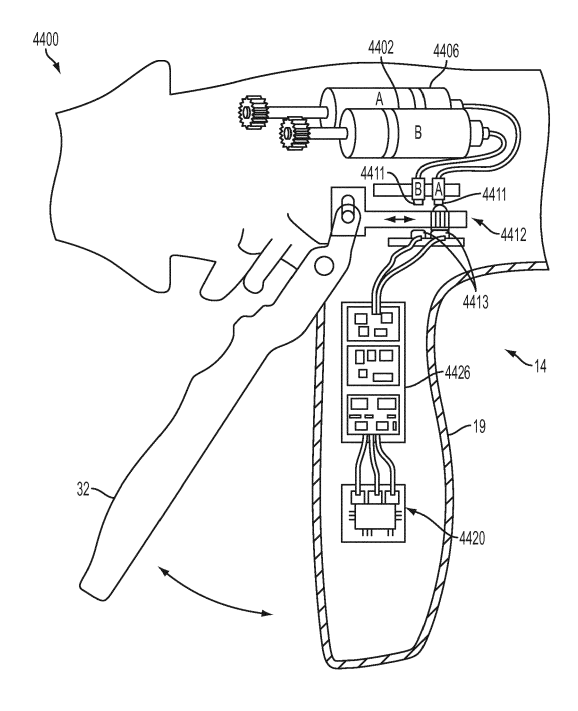
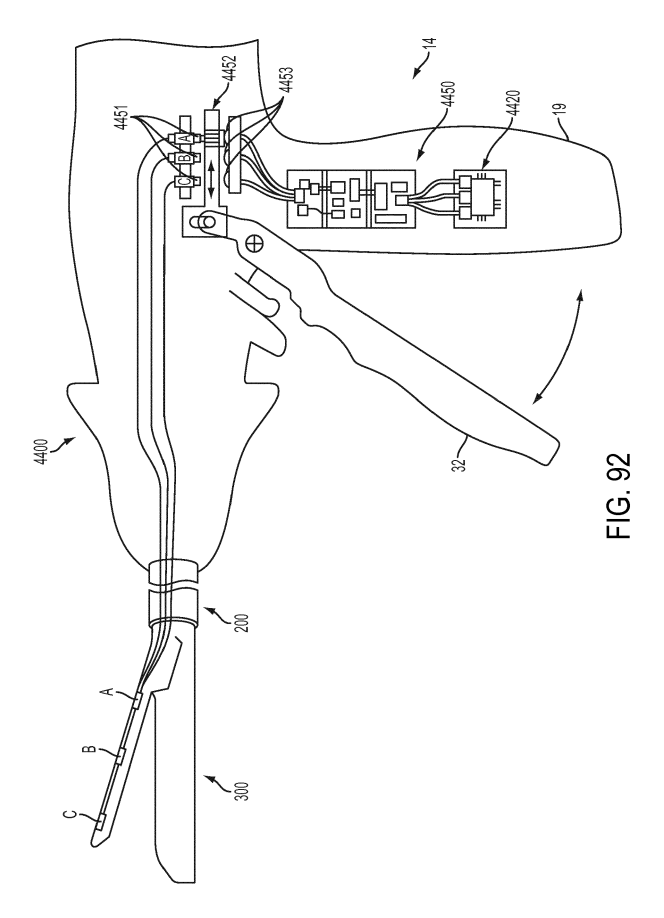
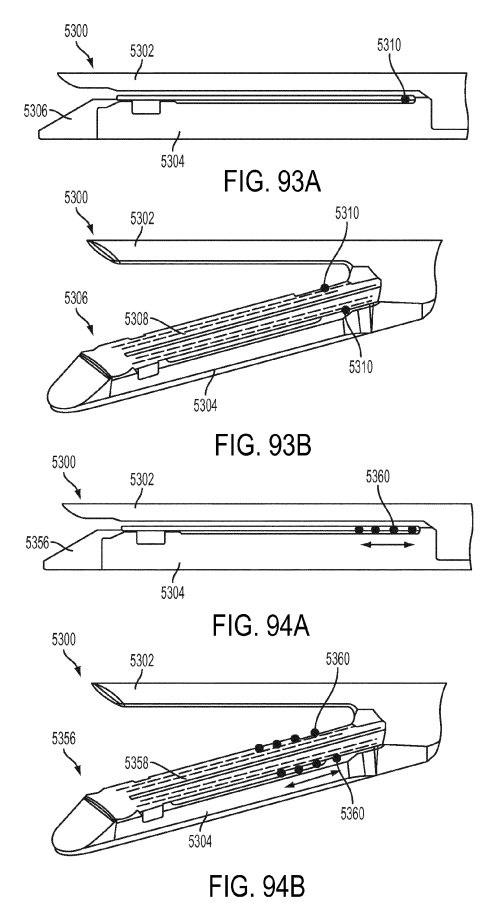


FIG. 91







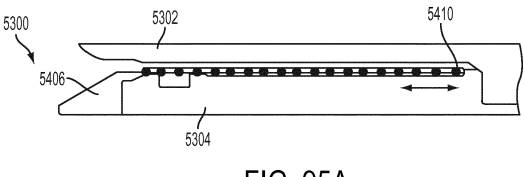


FIG. 95A

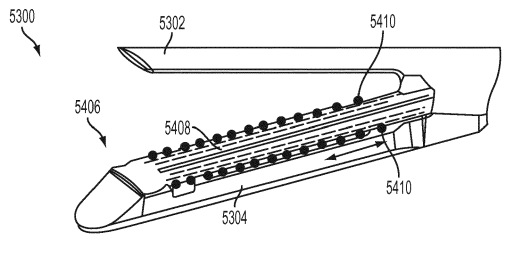
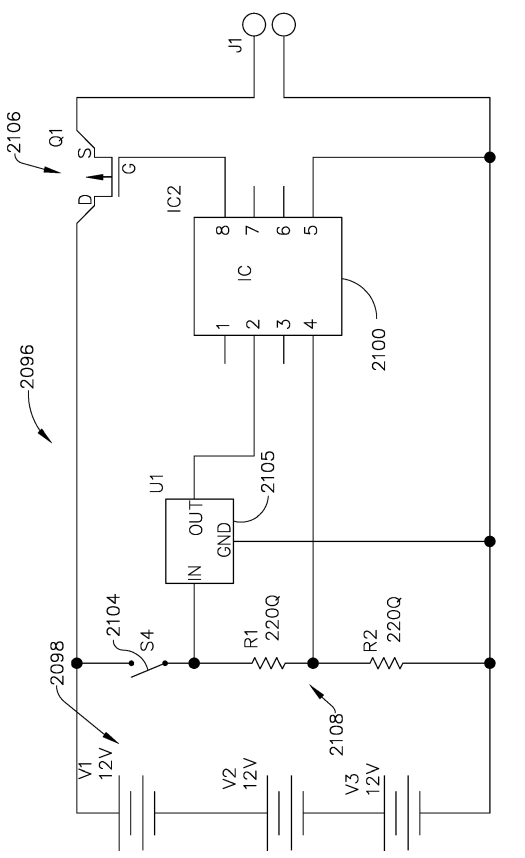
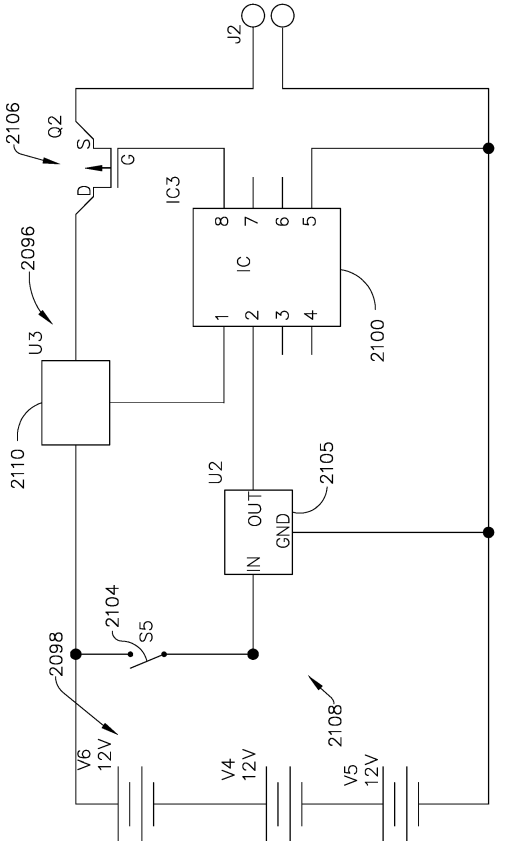
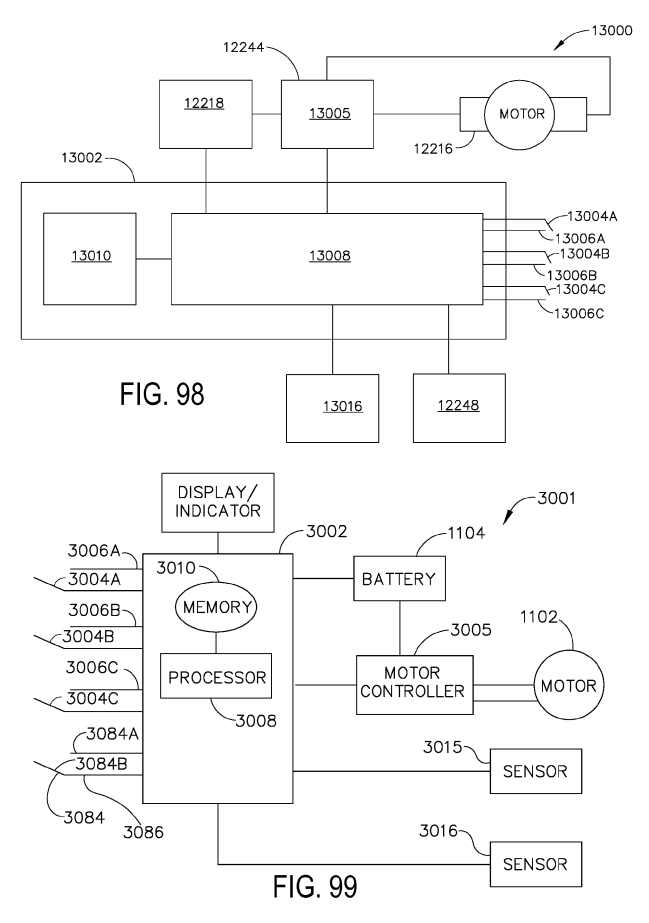


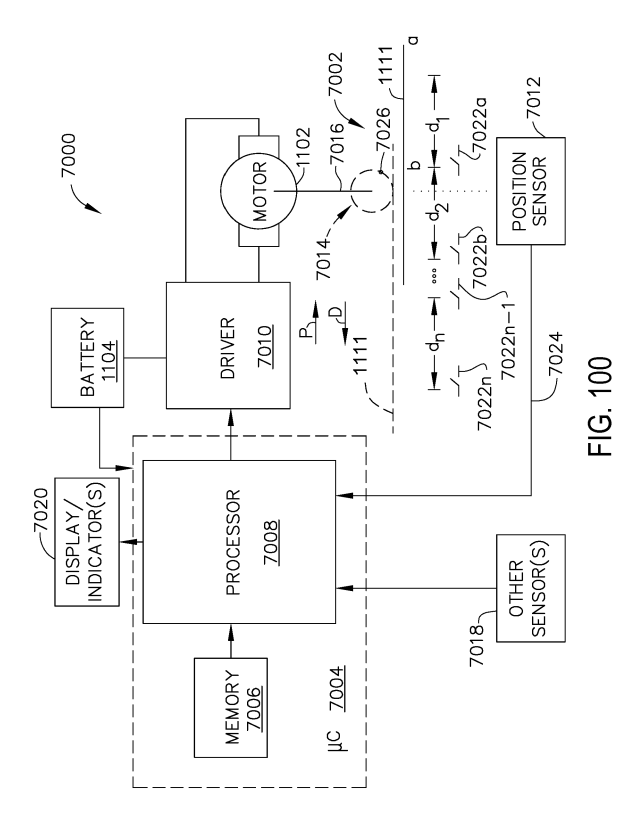
FIG. 95B

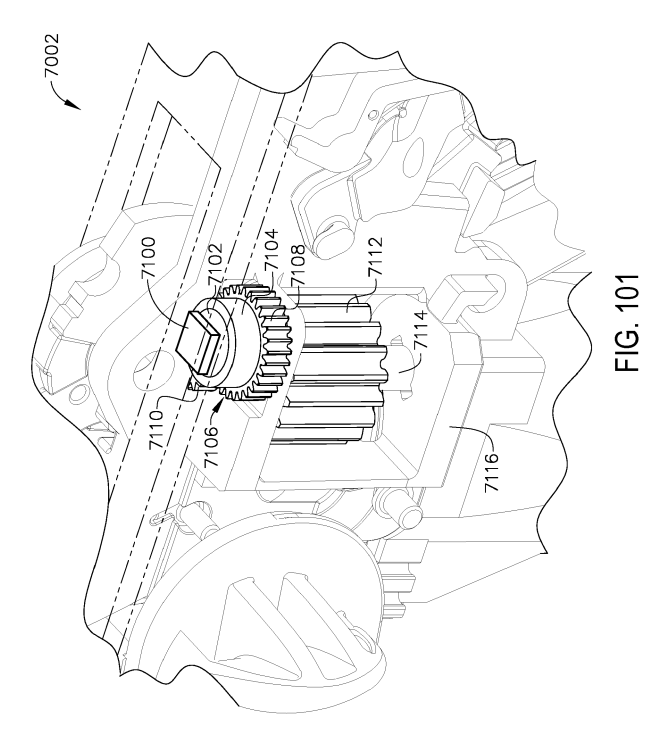


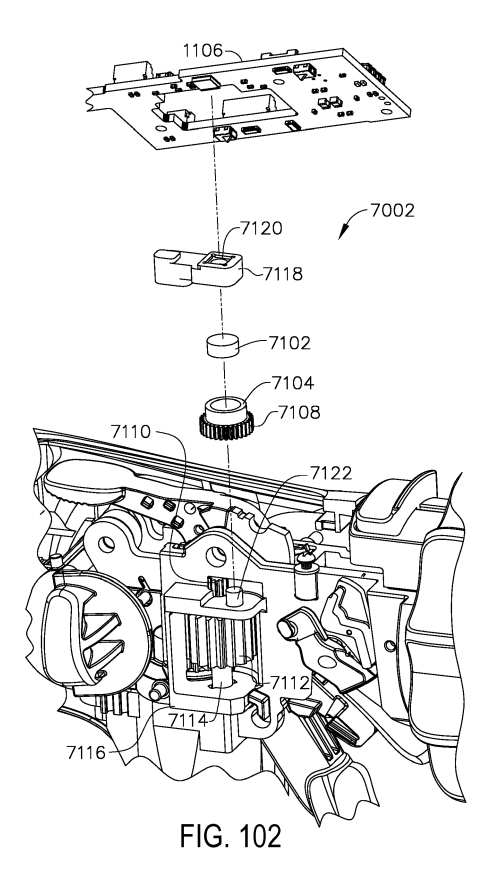


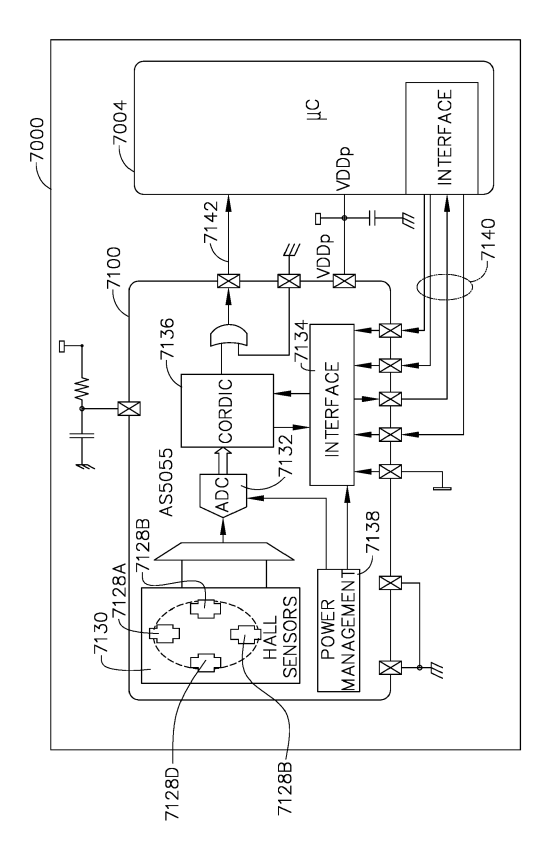


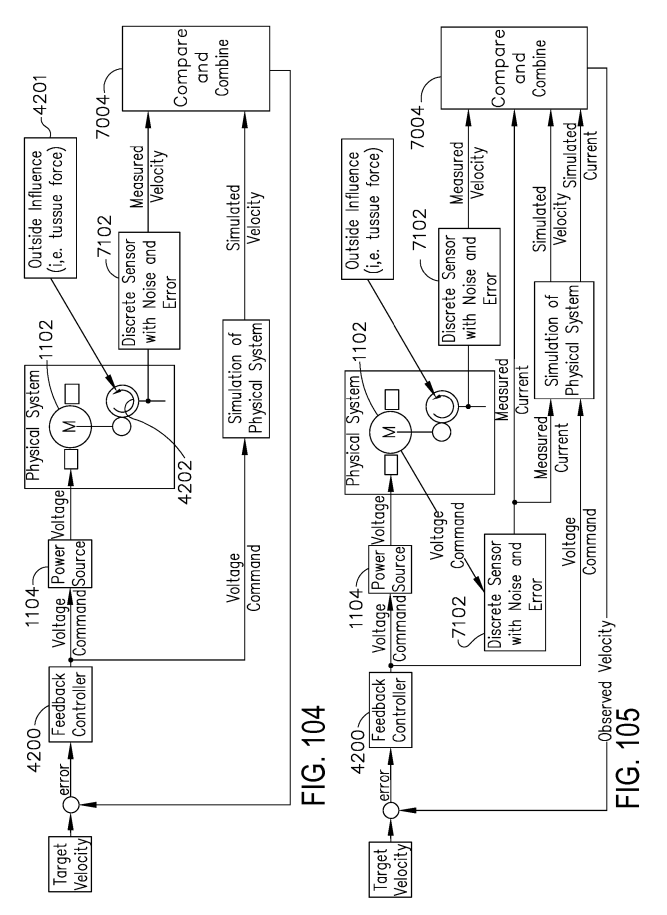












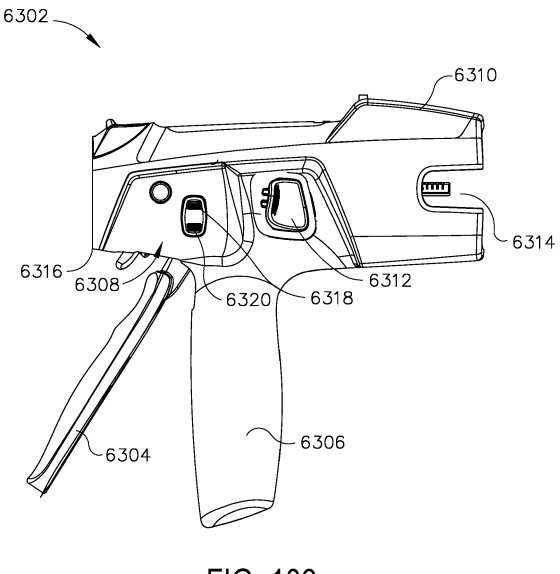
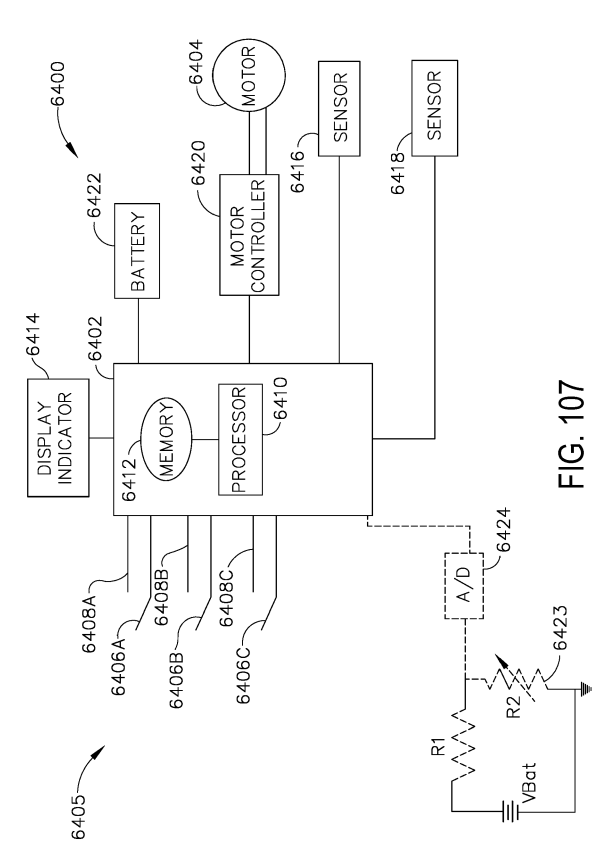
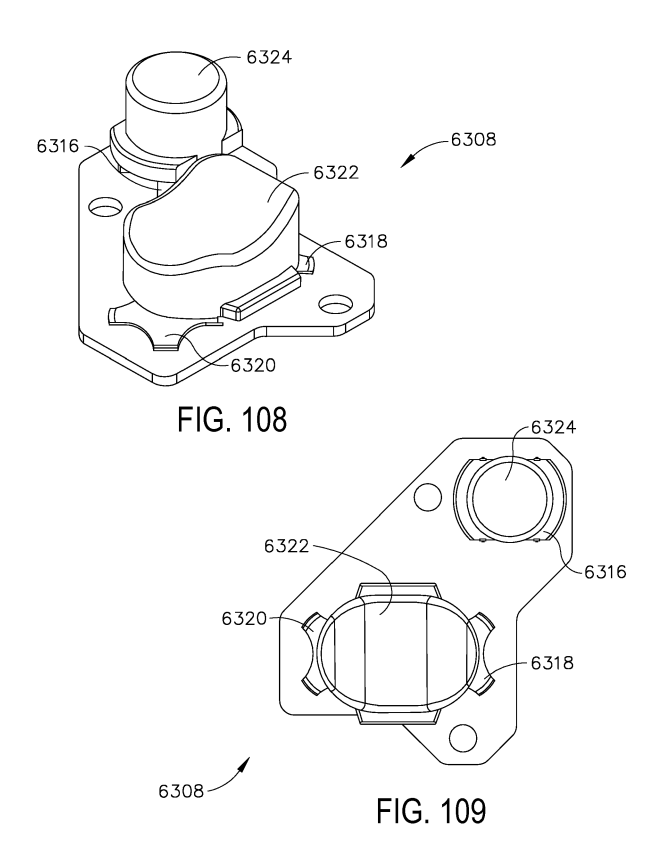
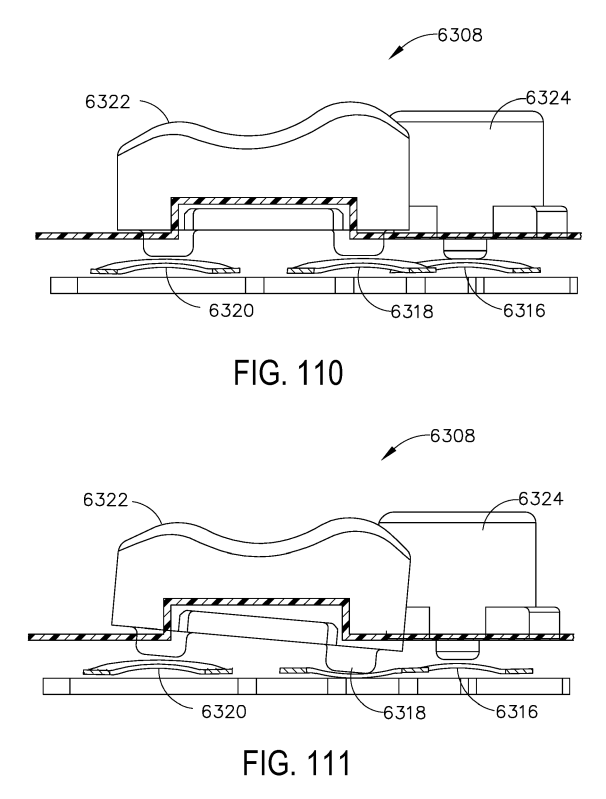
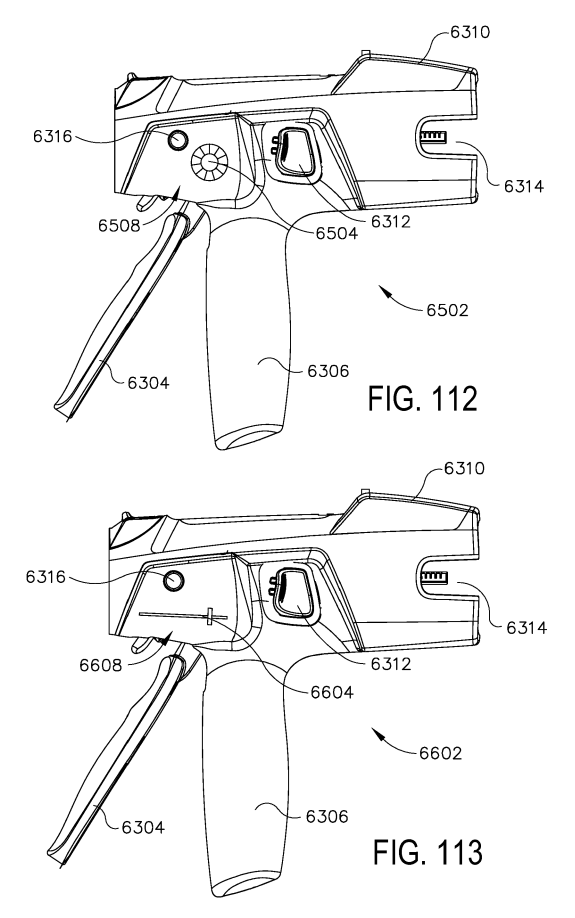


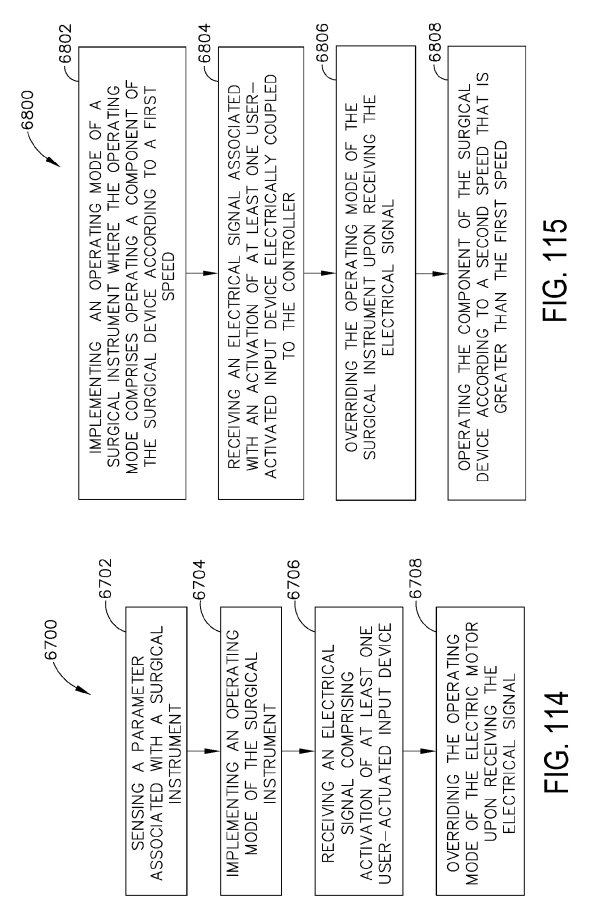
FIG. 106















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Application Number EP 17 16 6596

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	X	US 2012/211542 A1 (23 August 2012 (20] * paragraph [0012]	(RACENET DAVID C [US]) 2-08-23) *	1	
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					TECHNICAL FIELDS SEARCHED (IPC)
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1		The present search report has been drawn up for all claims			
50		Place of search Munich	Date of completion of the search 29 August 2017	Hel	Examiner d, Günter
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55	A:teo O:noi P:inte	hnological background n-written disclosure rrmediate document	& : member of the sa document	r, corresponding	

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